526 Rec'd PCT/PTO 1 6 JUL 2001

ATTORNEY'S DOCKET NUMBER U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE EORM PTO-1390 TRANSMITTAL LETTER TO THE UNITED STATES PF-0662 USN DESIGNATED/ELECTED OFFICE (DO/EO/US) TO BE AS INDIAN OF SERVICE TO BE AS INDIAN OF SE CONCERNING A FILING UNDER 35 U.S.C. 371 PRIORITY DATE CLAIMED INTERNATIONAL APPLICATION NO INTERNATIONAL FILING DATE

PCT/US00/02237

28 January 2000

20

29 January 1999

TITLE OF INVENTION

CANCER-ASSOCIATED PROTEINS

APPLICANT(S) FOR DO/EO/US

INCYTE PHARMACEUTICALS, INC.; TANG, Y. Tom; LAL, Preeti; HILLMAN, Jennifer L.; YUE, Henry; AZIMZAI, Yalda; LU, Dyung Aina M.; BAUGHN, Mariah R.; TRAN, Bao; SHIH, Leo L.; AU-YOUNG, Janice

Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:

- This is the FIRST submission of items concerning a filing under 35 U.S.C. 371.
- This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371.
- 3.

 This is an express request to promptly begin national examination procedures (35 U.S.C. 371 (f)). 4. □ The US has been elected by the expiration of 19 months from the priority date (PCT Article 31).
- 5.

 A copy of the International Application as filed (35 U.S.C. 371(c)(2))
 - a. □ is attached hereto (required only if not communicated by the International Bureau)
 - b.

 has been communicated by the International Bureau.
- c. is not required, as the application was filed in the United States Receiving Office (RO/US). ☐ An English language translation of the International Application as filed (35 U.S.C. 371(c)(2)).
- 7.

 Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3))
- a. \square are attached hereto (required only if not communicated by the International Bureau).

 - b.

 have been communicated by the International Bureau.
 - c.

 have not been made; however, the time limit for making such amendments has NOT expired.
 - d.

 have not been made and will not be made.
- 8. ☐ An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)). 9.

 An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)). 10.□ An English language translation of the annexes to the International Preliminary Examination Report under

PCT Article 36 (35 U.S.C. 371(c)(5)). Items 11 to 16 below concern document(s) or information included:

- □ An Information Disclosure Statement under 37 CFR 1.97 and 1.98.
- 12. ☐ An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.27 and 3 31 is included.
- □ A FIRST preliminary amendment.
 - A SECOND or SUBSEQUENT preliminary amendment.
- □ A substitute specification.
- □ A change of power of attorney and/or address letter.
- 1) Transmittal Letter (2 pp, in duplicate)
- 2) Return Postcard
- 3) Express Mail Label No.: EL 856 154 129 US
- 4) Request to Transfer

JC18 Rec'd PCT/PTO 1 6 JUL 2001

U.S. APPLICACION NO 8 8 9 6 7 TR 6) INTERNATIONAL APPLICATION DE ASSIGNER DE TRUBONO 2237		ICATION NO	ON NO - ATTORNEY'S DOCKET NUMBER PF-0662 USN			
17. 8 The following fies are submitted BASIC NATIONAL FEE (37 CFR L492(a)(1)-(5): Neither international preliminary examination fee (37 CFR L482) nor internationalsearch fee (37 CFR L445(a)(2)) paid to USPTO and internationalsearch for four not prepared bythe EPO or JPO\$1000.00 Clinternational preliminary examination fee (37 CFR L482) not paid to USPTO but international Search Report prepared bythe EPO or JPO\$600.00 International preliminary examination fee (37 CFR L482) not paid to USPTO but international search fee (37 CFR L482)) paid to USPTO\$710.00 Siluternational preliminary examination fee paid to USPTO 37 CFR L482) but all claims dld not satisfy provisions of PCT Article 33(1)-(4)5090.00 Clinternational preliminary examination fee paid to USPTO (37 CFR L482) and all elsims satisfied provisions of PCT Article 33(1)-(4)5100.00						
ENTER APPROPRIATE BASIC FEE AMOUNT =					\$690.00	
Surcharge of \$130.00 for furnishing the oath or declaration later than □ 20 □ 30 months from the earliest claimed priority date (37 CFR 1 492(e)).					s	
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE			
Total Claims	23 =	3	X \$ 18.00		\$ 54.00	
Independent Chims	2 =	0	X \$ 80.00		s	
MULTIPLE DEPEND	ENTCLAIM(S) (if applic	able)	+ \$270.00		s	
TOTAL OF ABOVE CALCULATIONS =					\$744.00	
□ Applicant clums small entity status. See 37 CFR 1.27. The fees indicated above are reduced by 1/2.					s	
SUBTOTAL =					\$744.00	
Processing fee of \$130.00 for furnishing the Englsh translation later than □ 20 □ 30 months from the earliest claimed priority date (37 CFR 1492(f)).					s	
TOTAL NATIONAL FEE =					\$744.00	
Fee for recording the enchsed assignment (37 CFR 1.21(h)). The assignment must be accompanied by the appropriate cover sheet (37 CFR 3.28, 3 31) \$40.00 per property +					s	
TOTAL FEES ENCLOSED =					\$744 00	
					Amount to be Refunded	s
					Charged	s
c. The Commission overpayment to	v Deposit Account No. 05 ter is hereby authorized to Deposit Account No. 09-C propriate time limit und to restore the application SPONDENCE TO	cover the above fees to 100 in the amount of \$\frac{5}{2}\$ charge anyadditional fees w 108. A dupleate copyof 1d or 37 CFR 1.494 or 1.4951 n to pending status. SIGRAFURE NAME: Diana Hand REGISTRATION NI	744 00	, a petition to	any	137(a) or (b)) must

1

PTO/PET RSC 11 6 JUL 2001

WO 00/44900

5

10

15

20

25

30

man from the first the fir

NUCLEIC-ACID BINDING PROTEINS

TECHNICAL FIELD

This invention relates to nucleic acid and amino acid sequences of nucleic-acid binding proteins and to the use of these sequences in the diagnosis, treatment, and prevention of reproductive, immune, and neurological disorders, and cell proliferative disorders including cancer.

BACKGROUND OF THE INVENTION

Multicellular organisms are comprised of diverse cell types that differ dramatically both in structure and function. The identity of a cell is determined by its characteristic pattern of gene expression, and different cell types express overlapping but distinct sets of genes throughout development. Spatial and temporal regulation of gene expression is critical for the control of cell proliferation, cell differentiation, apoptosis, and other processes that contribute to organismal development. Furthermore, gene expression is regulated in response to extracellular signals that mediate cell-cell communication and coordinate the activities of different cell types. Appropriate gene regulation also ensures that cells function efficiently by expressing only those genes whose functions are required at a given time.

Transcriptional regulatory proteins are essential for the control of gene expression. Some of these proteins function as transcription factors that initiate, activate, repress, or terminate gene transcription. Transcription factors generally bind to promoter, enhancer, or upstream regulatory regions of a gene in a sequence-specific manner, although some factors bind regulatory elements within or downstream of the coding region, Transcription factors may bind to a specific region of DNA singly or as a complex with other accessory factors. (Reviewed in Lewin, B. (1990) Genes IV, Oxford University Press, New York, NY, pp. 554-570.)

The double helix structure and repeated sequences of DNA create topological and chemical features which can be recognized by transcription factors. These features include hydrogen bond donor and acceptor groups, hydrophobic patches, major and minor grooves, and regular repeated stretches of sequence which induce distinct bends in the helix. Typically, transcription factors recognize specific DNA sequence motifs of about 20 nucleotides in length. Multiple adjacent transcription factor-binding motifs may be required for gene regulation.

Many transcription factors incorporate DNA-binding structural motifs which comprise either α helices or β sheets that bind to the major groove of DNA. Four well-characterized structural motifs are helix-turn-helix, zinc finger, leucine zipper, and helix-loop-helix. Proteins containing these motifs may act alone as monomers or form homo- or heterodimers that interact with DNA.

The helix-turn-helix motif consists of two α helices connected at a fixed angle by a short

1

chain of amino acids. One of the helices binds to the major groove. Helix-turn-helix motifs are exemplified by the homeobox motif which is present in homeodomain proteins. These proteins are critical for specifying the anterior-posterior body axis during development and are conserved throughout the animal kingdom. The Antennapedia and Ultrabithorax proteins of <u>Drosophila melanogaster</u> are prototypical homeodomain proteins. (Pabo, C.O. and R.T. Sauer (1992) Ann. Rev. Biochem. 61:1053-1095.)

The zinc finger motif, which binds zinc ions, generally contains tandem repeats of about 30 amino acids consisting of periodically spaced cysteine and histidine residues. Examples of this sequence pattern include the C2H2-type and the C3HC4-type zinc fingers, and the PHD domain. (Lewin, <u>supra</u>; Aasland, R., et al. (1995) Trends Biochem. Sci 20:56 - 59.) Zinc finger proteins each contain an α helix and an antiparallel β sheet whose proximity and conformation are maintained by the zinc ion. Contact with DNA is made by the arginine preceding the α helix and by the second, third, and sixth residues of the α helix. Variants of the zinc finger motif include poorly defined cysteine-rich motifs which bind zinc or other metal ions. These motifs may not contain histidine residues and are generally nonrepetitive.

The leucine zipper motif comprises a stretch of amino acids rich in leucine which can form an amphipathic α helix. This structure provides the basis for dimerization of two leucine zipper proteins. The region adjacent to the leucine zipper is usually basic, and upon protein dimerization, is optimally positioned for binding to the major groove. Proteins containing such motifs are generally referred to as bZIP transcription factors.

The helix-loop-helix motif (HLH) consists of a short α helix connected by a loop to a longer α helix. The loop is flexible and allows the two helices to fold back against each other and to bind to DNA. The transcription factor Myc contains a prototypical HLH motif.

20

25

Most transcription factors contain characteristic DNA binding motifs, and variations on the above motifs and new motifs have been and are currently being characterized. (Faisst, S. and S. Meyer (1992) Nucl. Acids Res. 20:3-26.)

Mutations in transcription factors contribute to oncogenesis. This is likely due to the role of transcription factors in the expression of genes involved in cell proliferation. For example, mutations in transcription factors encoded by proto-oncogenes, such as Fos, Jun, Myc, Rel, and Spil, may be oncogenic due to increased stimulation of cell proliferation. Conversely, mutations in transcription factors encoded by tumor suppressor genes, such as p53, RB1, and WT1, may be oncogenic due to decreased inhibition of cell proliferation. (Latchman, D. (1995) Gene Regulation: A Eukaryotic Perspective, Chapman and Hall, London, UK, pp 242-255.)

Gene expression is also affected by chromatin-associated proteins. In the nucleus, DNA is

Lines & with the english of the control of the cont

packaged into chromatin, the compact organization of which limits the accessibility of DNA to transcription factors and plays a key role in gene regulation. (Lewin, supra, pp. 409-410.) The compact structure of chromatin is determined and influenced by chromatin-associated proteins such as histones, high mobility group (HMG) proteins, helicases, and chromodomain proteins. There are five classes of histones, H1, H2A, H2B, H3, and H4, all of which are highly basic, low molecular weight proteins. The fundamental unit of chromatin, the nucleosome, consists of 200 base pairs of DNA associated with two copies each of H2A, H2B, H3, and H4. H1 links adjacent nucleosomes. HMG proteins are low molecular weight, non-histone proteins that may play a role in unwinding DNA and stabilizing single-stranded DNA. Helicases, which are DNA-dependent ATPases, unwind DNA, allowing access for transcription factors. Chromodomain proteins play a key role in the formation of highly-compacted, transcriptionally silent heterochromatin.

Much of the regulation of gene expression in eucaryotic cells occurs at the posttranscriptional level. Messenger RNAs (mRNA), which are produced in the cell nucleus from primary transcripts of protein-encoding genes, are processed and transported to the cytoplasm where the protein synthesis machinery is located. RNA-binding proteins are a group of proteins that participate in the processing, editing, transport, localization, and posttranscriptional regulation of mRNAs, and comprise the protein component of ribosomes as well. The RNA-binding activity of many of these proteins is mediated by a series of RNA-binding motifs identified within them. These domains include the RNP motif, the arginine-rich motif, the RGG box, and the KH motif. (Reviewed in Burd, C. G. and Dreyfuss, G. (1994) Science 265:615 - 621.) The RNP motif is the most widely found and best characterized of these motifs. The RNP motif is composed of 90-100 amino acids which form an RNA-binding domain and is found in one or more copies in proteins that bind pre-mRNA, mRNA, pre-ribosomal RNA, and small nuclear RNAs. The RNP motif is composed of two short sequences (RNP-1 and RNP-2) and a number of other mostly hydrophobic, conserved amino acids interspersed throughout the motif. (Burd, supra; ExPASy PROSITE document PDOC0030.)

15

25

Many neoplastic disorders in humans can be attributed to inappropriate gene expression.

Malignant cell growth may result from either excessive expression of tumor promoting genes or insufficient expression of tumor suppressor genes. (Cleary, M.L. (1992) Cancer Surv. 15:89-104.)

Chromosomal translocations may also produce chimeric loci which fuse the coding sequence of one gene with the regulatory regions of a second unrelated gene. Such an arrangement often results in inappropriate gene transcription. The Wilms tumor suppressor gene product, WT1, is a protein containing a DNA-binding domain consisting of four zinc fingers and a proline-glutamine rich region capable of regulating transcription. (ExPASy PROSITE document PR00049.) Deletions of the WT1 gene, or point mutations which destroy the DNA-binding activity of the protein are associated with development of the pediatric nephroblastoma, Wilms tumor, and Denys-Drash syndrome. (Rauscher,

20

30

PCT/US00/02237

F.J. (1993) FASEB J. 7:896-903.)

Certain proteins enriched in glutamine are associated with various neurological disorders including spinocerebellar ataxia, bipolar effective disorder, schizophrenia, and autism. (Margolis, R.L. et al. (1997) Human Genetics 100:114-122.) These proteins contain regions with as many as 15 or more consecutive glutamine residues and may function as transcription factors with a potential role in regulation of neurodevelopment or neuroplasticity.

The immune system responds to infection or trauma by activating a cascade of events that coordinate the progressive selection, amplification, and mobilization of cellular defense mechanisms. A complex and balanced program of gene activation and repression is involved in this process.

However, hyperactivity of the immune system as a result of improper or insufficient regulation of gene expression may result in considerable tissue or organ damage. This damage is well documented in immunological responses associated with arthritis, allergens, heart attack, stroke, and infections. (Harrison's Principles of Internal Medicine, 13/e, McGraw Hill, Inc. and Teton Data Systems Software, 1996.) In particular, a zinc finger protein termed Staf50 (for Stimulated trans-acting factor of 50 kDa) is a transcriptional regulator and is induced in various cell lines by interferon-I and -II. Staf50 appears to mediate the antiviral activity of interferon by down-regulating the viral transcription directed by the long terminal repeat promoter region of human immunodeficiency virus type-1 in transfected cells. (Tissot, C. (1995) J. Biol. Chem. 270:14891-14898.)

Furthermore, the generation of multicellular organisms is based upon the induction and coordination of cell differentiation at the appropriate stages of development. Central to this process is differential gene expression, which confers the distinct identities of cells and tissues throughout the body. Failure to regulate gene expression during development could result in developmental disorders.

The discovery of new nucleic-acid binding proteins and the polynucleotides encoding them satisfies a need in the art by providing new compositions which are useful in the diagnosis, prevention, and treatment of reproductive, immune, and neurological disorders, and cell proliferative disorders including cancer.

SUMMARY OF THE INVENTION

The invention features purified polypeptides, protnames, referred to collectively as "ABBR" and individually as "NuABP-1," "NuABP-2," "NuABP-3," "NuABP-4," "NuABP-5," "NuABP-6," "NuABP-7," "NuABP-8," "NuABP-9," "NuABP-10" "NuABP-11," "NuABP-12," "NuABP-13," "NuABP-14," "NuABP-15," "NuABP-16," "NuABP-17," "NuABP-18," "NuABP-19," "NuABP-20," "NuABP-21," "NuABP-22," "NuABP-23," "NuABP-24," "NuABP-25," "NuABP-26," "NuABP-27," "NuABP-28," "NuABP-33," "NuABP-31," "NuABP-32," "NuABP-33,"

"NuABP-34," "NuABP-35," "NuABP-36," "NuABP-37," "NuABP-38," "NuABP-39." "NuABP-40" "NuABP-41," "NuABP-42," "NuABP-43," "NuABP-44," "NuABP-45," "NuABP-46," "NuABP-47," "NuABP-48," "NuABP-49," "NuABP-50" "NuABP-51," "NuABP-52," "NuABP-53," "NuABP-54," and "NuABP-55." In one aspect, the invention provides an isolated polypeptide comprising a) an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, or d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55. In one alternative, the invention provides an isolated polypeptide comprising the amino acid sequence of SEO ID NO:1-55.

The invention further provides an isolated polynucleotide encoding a polypeptide comprising a) an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, c) a biologically active fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, or d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55. In one alternative, the polynucleotide is selected from the group consisting of SEQ ID NO:56-110.

10

20

25

30

35

Additionally, the invention provides a recombinant polynucleotide comprising a promoter sequence operably linked to a polynucleotide encoding a polypeptide comprising a) an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, c) a biologically active fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, or d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55. In one alternative, the invention provides a cell transformed with the recombinant polynucleotide. In another alternative, the invention provides a transgenic organism comprising the recombinant polynucleotide.

The invention also provides a method for producing a polypeptide comprising a) an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, c) a biologically active fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, or d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55. The method comprises a) culturing a cell under conditions suitable for expression of the polypeptide, wherein said cell is transformed with a recombinant polynucleotide comprising a promoter sequence operably linked to a polynucleotide encoding the polypeptide, and b) recovering the polypeptide so expressed.

Additionally, the invention provides an isolated antibody which specifically binds to a polypeptide comprising a) an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, c) a biologically active fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, or d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55

The invention further provides an isolated polynucleotide comprising a) a polynucleotide sequence selected from the group consisting of SEQ ID NO:56-110, b) a naturally occurring polynucleotide sequence having at least 90% sequence identity to a polynucleotide sequence selected from the group consisting of SEQ ID NO:56-110, c) a polynucleotide sequence complementary to a), or d) a polynucleotide sequence complementary to b). In one alternative, the polynucleotide comprises at least 60 contiguous nucleotides.

10

15

20

25

30

35

Additionally, the invention provides a method for detecting a target polynucleotide in a sample, said target polynucleotide having a sequence of a polynucleotide comprising a) a polynucleotide sequence selected from the group consisting of SEQ ID NO:56-110, b) a naturally occurring polynucleotide sequence having at least 90% sequence identity to a polynucleotide sequence complementary to a), or d) a polynucleotide sequence complementary to an experimentary to said target polynucleotide in the sample, and which probe specifically hybridizes to said target polynucleotide, under conditions whereby a hybridization complex is formed between said probe and said target polynucleotide, and b) detecting the presence or absence of said hybridization complex, and optionally, if present, the amount thereof. In one alternative, the probe comprises at least 30 contiguous nucleotides. In another alternative, the probe comprises at least 60 contiguous nucleotides.

The invention further provides a pharmaceutical composition comprising an effective amount of a polypeptide comprising a) an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, c) a biologically active fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, or d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, and a pharmaceutically acceptable excipient. The invention additionally provides a method of treating a disease or condition associated with decreased expression of functional NuABP, comprising administering to a patient in need of such treatment the pharmaceutical composition.

The invention also provides a method for screening a compound for effectiveness as an agonist of a polypeptide comprising a) an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, c) a biologically sactive fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, or d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55. The method comprises a) exposing a sample comprising the polypeptide to a compound, and b) detecting agonist activity in the sample. In one alternative, the invention provides a pharmaceutical composition comprising an agonist compound identified by the method and a pharmaceutically acceptable excipient. In another alternative, the invention provides a method of treating a disease or condition associated with decreased expression of functional NuABP, comprising administering to a patient in need of such treatment the pharmaceutical composition.

Additionally, the invention provides a method for screening a compound for effectiveness as an antagonist of a polypeptide comprising a) an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, c) a biologically active fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55, or d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1-55. The method comprises a) exposing a sample comprising the polypeptide to a compound, and b) detecting antagonist activity in the sample. In one alternative, the invention provides a pharmaceutical composition comprising an antagonist compound identified by the method and a pharmaceutically acceptable excipient. In another alternative, the invention provides a method of treating a disease or condition associated with overexpression of functional NuABP, comprising administering to a patient in need of such treatment the pharmaceutical composition.

The invention further provides a method for screening a compound for effectiveness in altering expression of a target polynucleotide, wherein said target polynucleotide comprises a sequence selected from the group consisting of SEQ ID NO:56-110, the method comprising a) exposing a sample comprising the target polynucleotide to a compound, and b) detecting altered expression of the target polynucleotide.

25

BRIEF DESCRIPTION OF THE TABLES

Table 1 shows polypeptide and nucleotide sequence identification numbers (SEQ ID NOs), clone identification numbers (clone IDs), cDNA libraries, and cDNA fragments used to assemble full-length sequences encoding NuABP.

Table 2 shows features of each polypeptide sequence, including potential motifs, homologous sequences, and methods, algorithms, and searchable databases used for analysis of NuABP.

Table 3 shows selected fragments of each nucleic acid sequence: the tissue-specific expression patterns of each nucleic acid sequence as determined by northern analysis; diseases, disorders, or conditions associated with these tissues; and the vector into which each cDNA was cloned.

Table 4 describes the tissues used to construct the cDNA libraries from which cDNA clones encoding NuABP were isolated.

Table 5 shows the tools, programs, and algorithms used to analyze NuABP, along with applicable descriptions, references, and threshold parameters.

DESCRIPTION OF THE INVENTION

Before the present proteins, nucleotide sequences, and methods are described, it is understood that this invention is not limited to the particular machines, materials and methods described, as these may vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to limit the scope of the present invention which will be limited only by the appended claims.

It must be noted that as used herein and in the appended claims, the singular forms "a," "an," and "the" include plural reference unless the context clearly dictates otherwise. Thus, for example, a reference to "a host cell" includes a plurality of such host cells, and a reference to "an antibody" is a reference to one or more antibodies and equivalents thereof known to those skilled in the art, and so forth.

Unless defined otherwise, all technical and scientific terms used herein have the same meanings as commonly understood by one of ordinary skill in the art to which this invention belongs. Although any machines, materials, and methods similar or equivalent to those described herein can be used to practice or test the present invention, the preferred machines, materials and methods are now described. All publications mentioned herein are cited for the purpose of describing and disclosing the cell lines, protocols, reagents and vectors which are reported in the publications and which might be used in connection with the invention. Nothing herein is to be construed as an admission that the invention is not entitled to antedate such disclosure by virtue of prior invention.

DEFINITIONS

15

20

30

35

"NuABP" refers to the amino acid sequences of substantially purified NuABP obtained from any species, particularly a mammalian species, including bovine, ovine, porcine, murine, equine, and human, and from any source, whether natural, synthetic, semi-synthetic, or recombinant.

The term "agonist" refers to a molecule which intensifies or mimics the biological activity of

NuABP. Agonists may include proteins, nucleic acids, carbohydrates, small molecules, or any other compound or composition which modulates the activity of NuABP either by directly interacting with NuABP or by acting on components of the biological pathway in which NuABP participates.

An "allelic variant" is an alternative form of the gene encoding NuABP. Allelic variants may result from at least one mutation in the nucleic acid sequence and may result in altered mRNAs or in polypeptides whose structure or function may or may not be altered. A gene may have none, one, or many allelic variants of its naturally occurring form. Common mutational changes which give rise to allelic variants are generally ascribed to natural deletions, additions, or substitutions of nucleotides. Each of these types of changes may occur alone, or in combination with the others, one or more times in a given sequence.

10

20

35

"Altered" nucleic acid sequences encoding NuABP include those sequences with deletions, insertions, or substitutions of different nucleotides, resulting in a polypeptide the same as NuABP or a polypeptide with at least one functional characteristic of NuABP. Included within this definition are polymorphisms which may or may not be readily detectable using a particular oligonucleotide probe of the polynucleotide encoding NuABP, and improper or unexpected hybridization to allelic variants. with a locus other than the normal chromosomal locus for the polynucleotide sequence encoding NuABP. The encoded protein may also be "altered," and may contain deletions, insertions, or substitutions of amino acid residues which produce a silent change and result in a functionally equivalent NuABP. Deliberate amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues, as long as the biological or immunological activity of NuABP is retained. For example, negatively charged amino acids may include aspartic acid and glutamic acid, and positively charged amino acids may include lysine and arginine. Amino acids with uncharged polar side chains having similar hydrophilicity values may include: asparagine and glutamine; and serine and threonine. Amino acids with uncharged side chains having similar hydrophilicity values may include: leucine, isoleucine, and valine; glycine and alanine; and phenylalanine and tyrosine.

The terms "amino acid" and "amino acid sequence" refer to an oligopeptide, peptide, polypeptide, or protein sequence, or a fragment of any of these, and to naturally occurring or synthetic molecules. Where "amino acid sequence" is recited to refer to an amino acid sequence of a naturally occurring protein molecule, "amino acid sequence" and like terms are not meant to limit the amino acid sequence to the complete native amino acid sequence associated with the recited protein molecule.

"Amplification" relates to the production of additional copies of a nucleic acid sequence.

Amplification is generally carried out using polymerase chain reaction (PCR) technologies well known in the art.

The term "antagonist" refers to a molecule which inhibits or attenuates the biological activity of NuABP. Antagonists may include proteins such as antibodies, nucleic acids, carbohydrates, small molecules, or any other compound or composition which modulates the activity of NuABP either by directly interacting with NuABP or by acting on components of the biological pathway in which NuABP participates.

The term "antibody" refers to intact immunoglobulin molecules as well as to fragments thereof, such as Fab, F(ab')₂, and Fv fragments, which are capable of binding an epitopic determinant. Antibodies that bind NuABP polypeptides can be prepared using intact polypeptides or using fragments containing small peptides of interest as the immunizing antigen. The polypeptide or oligopeptide used to immunize an animal (e.g., a mouse, a rat, or a rabbit) can be derived from the translation of RNA, or synthesized chemically, and can be conjugated to a carrier protein if desired. Commonly used carriers that are chemically coupled to peptides include bovine serum albumin, thyroglobulin, and keyhole limpet hemocyanin (KLH). The coupled peptide is then used to immunize the animal.

The term "antigenic determinant" refers to that region of a molecule (i.e., an epitope) that makes contact with a particular antibody. When a protein or a fragment of a protein is used to immunize a host animal, numerous regions of the protein may induce the production of antibodies which bind specifically to antigenic determinants (particular regions or three-dimensional structures on the protein). An antigenic determinant may compete with the intact antigen (i.e., the immunogen used to elicit the immune response) for binding to an antibody.

15

20

25

35

The term "antisense" refers to any composition containing a nucleic acid sequence which is complementary to the "sense" strand of a specific nucleic acid sequence. Antisense molecules may be produced by any method including synthesis or transcription. Once introduced into a cell, the complementary nucleotides combine with natural sequences produced by the cell to form duplexes and to block either transcription or translation. The designation "negative" or "minus" can refer to the antisense strand, and the designation "positive" or "plus" can refer to the sense strand.

The term "biologically active" refers to a protein having structural, regulatory, or biochemical functions of a naturally occurring molecule. Likewise, "immunologically active" refers to the capability of the natural, recombinant, or synthetic NuABP, or of any oligopeptide thereof, to induce a specific immune response in appropriate animals or cells and to bind with specific antibodies.

The terms "complementary" and "complementarity" refer to the natural binding of polynucleotides by base pairing. For example, the sequence "5' A-G-T 3" bonds to the complementary sequence "3' T-C-A 5'." Complementarity between two single-stranded molecules may be "partial," such that only some of the nucleic acids bind, or it may be "complete," such that total complementarity exists between the single stranded molecules. The degree of complementarity

between nucleic acid strands has significant effects on the efficiency and strength of the hybridization between the nucleic acid strands. This is of particular importance in amplification reactions, which depend upon binding between nucleic acid strands, and in the design and use of peptide nucleic acid (PNA) molecules.

5

15

20

A "composition comprising a given polynucleotide sequence" and a "composition comprising a given amino acid sequence" refer broadly to any composition containing the given polynucleotide or amino acid sequence. The composition may comprise a dry formulation or an aqueous solution. Compositions comprising polynucleotide sequences encoding NuABP or fragments of NuABP may be employed as hybridization probes. The probes may be stored in freeze-dried form and may be associated with a stabilizing agent such as a carbohydrate. In hybridizations, the probe may be deployed in an aqueous solution containing salts (e.g., NaCl), detergents (e.g., sodium dodecyl sulfate; SDS), and other components (e.g., Denhardt's solution, dry milk, salmon sperm DNA. etc.).

"Consensus sequence" refers to a nucleic acid sequence which has been resequenced to resolve uncalled bases, extended using the XL-PCR kit (Perkin-Elmer, Norwalk CT) in the 5' and/or the 3' direction, and resequenced, or which has been assembled from the overlapping sequences of one or more Incyte Clones and, in some cases, one or more public domain ESTs, using a computer program for fragment assembly, such as the GELVIEW fragment assembly system (GCG, Madison WI). Some sequences have been both extended and assembled to produce the consensus sequence.

"Conservative amino acid substitutions" are those substitutions that, when made, least interfere with the properties of the original protein, i.e., the structure and especially the function of the protein is conserved and not significantly changed by such substitutions. The table below shows amino acids which may be substituted for an original amino acid in a protein and which are regarded as conservative amino acid substitutions.

	Original Residue	Conservative Substitution
25	Ala	Gly, Ser
	Arg	His, Lys
	Asn	Asp, Gln, His
	Asp	Asn, Glu
	Cys	Ala, Ser
30	Gln	Asn, Glu, His
	Glu	Asp, Gln, His
	Gly	Ala
	His	Asn, Arg, Gln, Glu
	Ile	Leu, Val
35	Leu	Ile, Val
	Lys	Arg, Gln, Glu
	Met	Leu, He
	Phe	His, Met, Leu, Trp, Tyr
	Ser	Cys, Thr
40	Thr	Ser, Val

Trp	Phe, Tyr
Tyr	His, Phe, Trp
Val	lle. Leu. Thr

Conservative amino acid substitutions generally maintain (a) the structure of the polypeptide backbone in the area of the substitution, for example, as a beta sheet or alpha helical conformation, (b) the charge or hydrophobicity of the molecule at the site of the substitution, and/or (c) the bulk of the side chain.

5

15

25

35

A "deletion" refers to a change in the amino acid or nucleotide sequence that results in the absence of one or more amino acid residues or nucleotides.

The term "derivative" refers to the chemical modification of a polypeptide sequence, or a polynucleotide sequence. Chemical modifications of a polynucleotide sequence can include, for example, replacement of hydrogen by an alkyl, acyl, hydroxyl, or amino group. A derivative polynucleotide encodes a polypeptide which retains at least one biological or immunological function of the natural molecule. A derivative polypeptide is one modified by glycosylation, pegylation, or any similar process that retains at least one biological or immunological function of the polypeptide from which it was derived.

A "fragment" is a unique portion of NuABP or the polynucleotide encoding NuABP which is identical in sequence to but shorter in length than the parent sequence. A fragment may comprise up to the entire length of the defined sequence, minus one nucleotide/amino acid residue. For example, a fragment may comprise from 5 to 1000 contiguous nucleotides or amino acid residues. A fragment used as a probe, primer, antigen, therapeutic molecule, or for other purposes, may be at least 5, 10, 15, 20, 25, 30, 40, 50, 60, 75, 100, 150, 250 or at least 500 contiguous nucleotides or amino acid residues in length. Fragments may be preferentially selected from certain regions of a molecule. For example, a polypeptide fragment may comprise a certain length of contiguous amino acids selected from the first 250 or 500 amino acids (or first 25% or 50% of a polypeptide) as shown in a certain defined sequence. Clearly these lengths are exemplary, and any length that is supported by the specification, including the Sequence Listing, tables, and figures, may be encompassed by the present embodiments.

A fragment of SEQ ID NO:56-110 comprises a region of unique polynucleotide sequence that specifically identifies SEQ ID NO:56-110, for example, as distinct from any other sequence in the same genome. A fragment of SEQ ID NO:56-110 is useful, for example, in hybridization and amplification technologies and in analogous methods that distinguish SEQ ID NO:56-110 from related polynucleotide sequences. The precise length of a fragment of SEQ ID NO:56-110 and the region of SEQ ID NO:56-110 to which the fragment corresponds are routinely determinable by one of ordinary skill in the art based on the intended purpose for the fragment.

A fragment of SEQ ID NO:1-55 is encoded by a fragment of SEO ID NO:56-110. A

fragment of SEQ ID NO:1-55 comprises a region of unique amino acid sequence that specifically identifies SEQ ID NO:1-55. For example, a fragment of SEQ ID NO:1-55 is useful as an immunogenic peptide for the development of antibodies that specifically recognize SEQ ID NO:1-55. The precise length of a fragment of SEQ ID NO:1-55 and the region of SEQ ID NO:1-55 to which the fragment corresponds are routinely determinable by one of ordinary skill in the art based on the intended purpose for the fragment.

The term "similarity" refers to a degree of complementarity. There may be partial similarity or complete similarity. The word "identity" may substitute for the word "similarity." A partially complementary sequence that at least partially inhibits an identical sequence from hybridizing to a target nucleic acid is referred to as "substantially similar." The inhibition of hybridization of the completely complementary sequence to the target sequence may be examined using a hybridization assay (Southern or northern blot, solution hybridization, and the like) under conditions of reduced stringency. A substantially similar sequence or hybridization probe will compete for and inhibit the binding of a completely similar (identical) sequence to the target sequence under conditions of reduced stringency. This is not to say that conditions of reduced stringency are such that non-specific binding is permitted, as reduced stringency conditions require that the binding of two sequences to one another be a specific (i.e., a selective) interaction. The absence of non-specific binding may be tested by the use of a second target sequence which lacks even a partial degree of complementarity (e.g., less than about 30% similarity or identity). In the absence of non-specific binding, the substantially similar sequence or probe will not hybridize to the second non-complementary target sequence.

10

20

35

The phrases "percent identity" and "% identity," as applied to polynucleotide sequences, refer to the percentage of residue matches between at least two polynucleotide sequences aligned using a standardized algorithm. Such an algorithm may insert, in a standardized and reproducible way, gaps in the sequences being compared in order to optimize alignment between two sequences, and therefore achieve a more meaningful comparison of the two sequences.

Percent identity between polynucleotide sequences may be determined using the default parameters of the CLUSTAL V algorithm as incorporated into the MEGALIGN version 3.12e sequence alignment program. This program is part of the LASERGENE software package, a suite of molecular biological analysis programs (DNASTAR, Madison WI). CLUSTAL V is described in Higgins, D.G. and P.M. Sharp (1989) CABIOS 5:151-153 and in Higgins, D.G. et al. (1992) CABIOS 8:189-191. For pairwise alignments of polynucleotide sequences, the default parameters are set as follows: Ktuple=2, gap penalty=5, window=4, and "diagonals saved"=4. The "weighted" residue weight table is selected as the default. Percent identity is reported by CLUSTAL V as the "percent similarity" between aligned polynucleotide sequence pairs.

Alternatively, a suite of commonly used and freely available sequence comparison algorithms is provided by the National Center for Biotechnology Information (NCBI) Basic Local Alignment Search Tool (BLAST) (Altschul, S.F. et al. (1990) J. Mol. Biol. 215:403-410), which is available from several sources, including the NCBI, Bethesda, MD, and on the Internet at

5 http://www.ncbi.nlm.nih.gov/BLAST/. The BLAST software suite includes various sequence analysis programs including "blastn," that is used to align a known polynucleotide sequence with other polynucleotide sequences from a variety of databases. Also available is a tool called "BLAST 2 Sequences" that is used for direct pairwise comparison of two nucleotide sequences. "BLAST 2 Sequences" can be accessed and used interactively at http://www.ncbi.nlm.nih.gov/gorf/bl2.html. The "BLAST 2 Sequences" tool can be used for both blastn and blastp (discussed below). BLAST programs are commonly used with gap and other parameters set to default settings. For example, to compare two nucleotide sequences, one may use blastn with the "BLAST 2 Sequences" tool Version

2.0.9 (May-07-1999) set at default parameters. Such default parameters may be, for example:

Matrix: BLOSUM62

15 Reward for match: I

Penalty for mismatch: -2

Open Gap: 5 and Extension Gap: 2 penalties

Gap x drop-off: 50

Expect: 10

Word Size: 11

Filter: on

20

25

30

35

Percent identity may be measured over the length of an entire defined sequence, for example, as defined by a particular SEQ ID number, or may be measured over a shorter length, for example, over the length of a fragment taken from a larger, defined sequence, for instance, a fragment of at least 20, at least 30, at least 40, at least 50, at least 70, at least 100, or at least 200 contiguous nucleotides. Such lengths are exemplary only, and it is understood that any fragment length supported by the sequences shown herein, in the tables, figures, or Sequence Listing, may be used to describe a length over which percentage identity may be measured.

Nucleic acid sequences that do not show a high degree of identity may nevertheless encode similar amino acid sequences due to the degeneracy of the genetic code. It is understood that changes in a nucleic acid sequence can be made using this degeneracy to produce multiple nucleic acid sequences that all encode substantially the same protein.

The phrases "percent identity" and "% identity," as applied to polypeptide sequences, refer to the percentage of residue matches between at least two polypeptide sequences aligned using a standardized algorithm. Methods of polypeptide sequence alignment are well-known. Some

alignment methods take into account conservative amino acid substitutions. Such conservative substitutions, explained in more detail above, generally preserve the hydrophobicity and acidity at the site of substitution, thus preserving the structure (and therefore function) of the polypeptide.

Percent identity between polypeptide sequences may be determined using the default

5 parameters of the CLUSTAL V algorithm as incorporated into the MEGALIGN version 3.12e
sequence alignment program (described and referenced above). For pairwise alignments of
polypeptide sequences using CLUSTAL V, the default parameters are set as follows: Ktuple=1, gap
penalty=3, window=5, and "diagonals saved"=5. The PAM250 matrix is selected as the default
residue weight table. As with polynucleotide alignments, the percent identity is reported by

10 CLUSTAL V as the "percent similarity" between aligned polypeptide sequence pairs.

Alternatively the NCBI BLAST software suite may be used. For example, for a pairwise comparison of two polypeptide sequences, one may use the "BLAST 2 Sequences" tool Version 2.0.9 (May-07-1999) with blastp set at default parameters. Such default parameters may be, for example:

Matrix: BLOSUM62

Open Gap: 11 and Extension Gap: 1 penalties

Gap x drop-off: 50

Expect: 10

15

20

25

30

Word Size: 3

Filter: on

Percent identity may be measured over the length of an entire defined polypeptide sequence, for example, as defined by a particular SEQ ID number, or may be measured over a shorter length, for example, over the length of a fragment taken from a larger, defined polypeptide sequence, for instance, a fragment of at least 15, at least 20, at least 30, at least 40, at least 50, at least 70 or at least 150 contiguous residues. Such lengths are exemplary only, and it is understood that any fragment length supported by the sequences shown herein, in the tables, figures or Sequence Listing, may be used to describe a length over which percentage identity may be measured.

"Human artificial chromosomes" (HACs) are linear microchromosomes which may contain DNA sequences of about 6 kb to 10 Mb in size, and which contain all of the elements required for stable mitotic chromosome segregation and maintenance.

The term "humanized antibody" refers to antibody molecules in which the amino acid sequence in the non-antigen binding regions has been altered so that the antibody more closely resembles a human antibody, and still retains its original binding ability.

"Hybridization" refers to the process by which a polynucleotide strand anneals with a complementary strand through base pairing under defined hybridization conditions. Specific hybridization is an indication that two nucleic acid sequences share a high degree of identity. Specific

hybridization complexes form under permissive annealing conditions and remain hybridized after the "washing" step(s). The washing step(s) is particularly important in determining the stringency of the hybridization process, with more stringent conditions allowing less non-specific binding, i.e., binding between pairs of nucleic acid strands that are not perfectly matched. Permissive conditions for annealing of nucleic acid sequences are routinely determinable by one of ordinary skill in the art and may be consistent among hybridization experiments, whereas wash conditions may be varied among experiments to achieve the desired stringency, and therefore hybridization specificity. Permissive annealing conditions occur, for example, at 68°C in the presence of about 6 x SSC, about 1% (w/v) SDS, and about 100 µg/ml denatured salmon sperm DNA.

Generally, stringency of hybridization is expressed, in part, with reference to the temperature under which the wash step is carried out. Generally, such wash temperatures are selected to be about 5°C to 20°C lower than the thermal melting point (T_m) for the specific sequence at a defined ionic strength and pH. The T_m is the temperature (under defined ionic strength and pH) at which 50% of the target sequence hybridizes to a perfectly matched probe. An equation for calculating T_m and conditions for nucleic acid hybridization are well known and can be found in Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual, 2nd ed., vol. 1-3, Cold Spring Harbor Press, Plainview NY; specifically see volume 2, chapter 9.

10

20

25

30

35

High stringency conditions for hybridization between polynucleotides of the present invention include wash conditions of 68°C in the presence of about 0.2 x SSC and about 0.1% SDS, for 1 hour. Alternatively, temperatures of about 65°C, 60°C, 55°C, or 42°C may be used. SSC concentration may be varied from about 0.1 to 2 x SSC, with SDS being present at about 0.1%. Typically, blocking reagents are used to block non-specific hybridization. Such blocking reagents include, for instance, denatured salmon sperm DNA at about 100-200 μ g/ml. Organic solvent, such as formamide at a concentration of about 35-50% v/v, may also be used under particular circumstances, such as for RNA:DNA hybridizations. Useful variations on these wash conditions will be readily apparent to those of ordinary skill in the art. Hybridization, particularly under high stringency conditions, may be suggestive of evolutionary similarity between the nucleotides. Such similarity is strongly indicative of a similar role for the nucleotides and their encoded polypeptides.

The term "hybridization complex" refers to a complex formed between two nucleic acid sequences by virtue of the formation of hydrogen bonds between complementary bases. A hybridization complex may be formed in solution (e.g., Cot or Rot analysis) or formed between one nucleic acid sequence present in solution and another nucleic acid sequence immobilized on a solid support (e.g., paper, membranes, filters, chips, pins or glass slides, or any other appropriate substrate to which cells or their nucleic acids have been fixed).

The words "insertion" and "addition" refer to changes in an amino acid or nucleotide

sequence resulting in the addition of one or more amino acid residues or nucleotides, respectively.

"Immune response" can refer to conditions associated with inflammation, trauma, immune disorders, or infectious or genetic disease, etc. These conditions can be characterized by expression of various factors, e.g., cytokines, chemokines, and other signaling molecules, which may affect cellular and systemic defense systems.

The term "microarray" refers to an arrangement of distinct polynucleotides on a substrate.

The terms "element" and "array element" in a microarray context, refer to hybridizable polynucleotides arranged on the surface of a substrate.

10

15

20

25

30

The term "modulate" refers to a change in the activity of NuABP. For example, modulation may cause an increase or a decrease in protein activity, binding characteristics. or any other biological, functional, or immunological properties of NuABP.

The phrases "nucleic acid" and "nucleic acid sequence" refer to a nucleotide, oligonucleotide, polynucleotide, or any fragment thereof. These phrases also refer to DNA or RNA of genomic or synthetic origin which may be single-stranded or double-stranded and may represent the sense or the antisense strand, to peptide nucleic acid (PNA), or to any DNA-like or RNA-like material.

"Operably linked" refers to the situation in which a first nucleic acid sequence is placed in a functional relationship with the second nucleic acid sequence. For instance, a promoter is operably linked to a coding sequence if the promoter affects the transcription or expression of the coding sequence. Generally, operably linked DNA sequences may be in close proximity or contiguous and, where necessary to join two protein coding regions, in the same reading frame.

"Peptide nucleic acid" (PNA) refers to an antisense molecule or anti-gene agent which comprises an oligonucleotide of at least about 5 nucleotides in length linked to a peptide backbone of amino acid residues ending in lysine. The terminal lysine confers solubility to the composition. PNAs preferentially bind complementary single stranded DNA or RNA and stop transcript elongation, and may be pegylated to extend their lifespan in the cell.

"Probe" refers to nucleic acid sequences encoding NuABP, their complements, or fragments thereof, which are used to detect identical, allelic or related nucleic acid sequences. Probes are isolated oligonucleotides or polynucleotides attached to a detectable label or reporter molecule. Typical labels include radioactive isotopes, ligands, chemiluminescent agents, and enzymes. "Primers" are short nucleic acids, usually DNA oligonucleotides, which may be annealed to a target polynucleotide by complementary base-pairing. The primer may then be extended along the target DNA strand by a DNA polymerase enzyme. Primer pairs can be used for amplification (and identification) of a nucleic acid sequence, e.g., by the polymerase chain reaction (PCR).

Probes and primers as used in the present invention typically comprise at least 15 contiguous

35 nucleotides of a known sequence. In order to enhance specificity, longer probes and primers may also

be employed, such as probes and primers that comprise at least 20, 25, 30, 40, 50, 60, 70, 80, 90, 100, or at least 150 consecutive nucleotides of the disclosed nucleic acid sequences. Probes and primers may be considerably longer than these examples, and it is understood that any length supported by the specification, including the tables, figures, and Sequence Listing, may be used.

Methods for preparing and using probes and primers are described in the references, for example Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual, 2nd ed., vol. 1-3, Cold Spring Harbor Press, Plainview NY; Ausubel et al., 1987, <u>Current Protocols in Molecular Biology</u>, Greene Publ. Assoc. & Wiley-Intersciences, New York NY; Innis et al., 1990, <u>PCR Protocols. A Guide to Methods and Applications</u>, Academic Press, San Diego CA. PCR primer pairs can be derived from a known sequence, for example, by using computer programs intended for that purpose such as Primer (Version 0.5, 1991, Whitehead Institute for Biomedical Research, Cambridge MA).

5

15

20

2.5

Oligonucleotides for use as primers are selected using software known in the art for such purpose. For example, OLIGO 4.06 software is useful for the selection of PCR primer pairs of up to 100 nucleotides each, and for the analysis of oligonucleotides and larger polynucleotides of up to 5,000 nucleotides from an input polynucleotide sequence of up to 32 kilobases. Similar primer selection programs have incorporated additional features for expanded capabilities. For example, the PrimOU primer selection program (available to the public from the Genome Center at University of Texas South West Medical Center, Dallas TX) is capable of choosing specific primers from megabase sequences and is thus useful for designing primers on a genome-wide scope. The Primer3 primer selection program (available to the public from the Whitehead Institute/MIT Center for Genome Research, Cambridge MA) allows the user to input a "mispriming library," in which sequences to avoid as primer binding sites are user-specified. Primer3 is useful, in particular, for the selection of oligonucleotides for microarrays. (The source code for the latter two primer selection programs may also be obtained from their respective sources and modified to meet the user's specific needs.) The PrimeGen program (available to the public from the UK Human Genome Mapping Project Resource Centre, Cambridge UK) designs primers based on multiple sequence alignments, thereby allowing selection of primers that hybridize to either the most conserved or least conserved regions of aligned nucleic acid sequences. Hence, this program is useful for identification of both unique and conserved oligonucleotides and polynucleotide fragments. The oligonucleotides and polynucleotide fragments identified by any of the above selection methods are useful in hybridization technologies, for example, as PCR or sequencing primers, microarray elements, or specific probes to identify fully or partially complementary polynucleotides in a sample of nucleic acids. Methods of oligonucleotide selection are not limited to those described above.

A "recombinant nucleic acid" is a sequence that is not naturally occurring or has a sequence
that is made by an artificial combination of two or more otherwise separated segments of sequence.

This artificial combination is often accomplished by chemical synthesis or, more commonly, by the artificial manipulation of isolated segments of nucleic acids, e.g., by genetic engineering techniques such as those described in Sambrook. supra. The term recombinant includes nucleic acids that have been altered solely by addition, substitution, or deletion of a portion of the nucleic acid. Frequently, a recombinant nucleic acid may include a nucleic acid sequence operably linked to a promoter sequence. Such a recombinant nucleic acid may be part of a vector that is used, for example, to transform a cell.

Alternatively, such recombinant nucleic acids may be part of a viral vector, e.g., based on a vaccinia virus, that could be use to vaccinate a mammal wherein the recombinant nucleic acid is expressed, inducing a protective immunological response in the mammal.

10

15

20

25

The term "sample" is used in its broadest sense. A sample suspected of containing nucleic acids encoding NuABP, or fragments thereof, or NuABP itself, may comprise a bodily fluid; an extract from a cell, chromosome, organelle, or membrane isolated from a cell; a cell; genomic DNA, RNA, or cDNA, in solution or bound to a substrate; a tissue; a tissue print; etc.

The terms "specific binding" and "specifically binding" refer to that interaction between a protein or peptide and an agonist, an antibody, an antagonist, a small molecule, or any natural or synthetic binding composition. The interaction is dependent upon the presence of a particular structure of the protein, e.g., the antigenic determinant or epitope, recognized by the binding molecule. For example, if an antibody is specific for epitope "A," the presence of a polypeptide containing the epitope A, or the presence of free unlabeled A, in a reaction containing free labeled A and the antibody will reduce the amount of labeled A that binds to the antibody.

The term "substantially purified" refers to nucleic acid or amino acid sequences that are removed from their natural environment and are isolated or separated, and are at least 60% free, preferably at least 75% free, and most preferably at least 90% free from other components with which they are naturally associated.

A "substitution" refers to the replacement of one or more amino acids or nucleotides by different amino acids or nucleotides, respectively.

"Substrate" refers to any suitable rigid or semi-rigid support including membranes, filters, chips, slides, wafers, fibers, magnetic or nonmagnetic beads, gels, tubing, plates, polymers, microparticles and capillaries. The substrate can have a variety of surface forms, such as wells, trenches, pins, channels and pores, to which polynucleotides or polypeptides are bound.

"Transformation" describes a process by which exogenous DNA enters and changes a recipient cell. Transformation may occur under natural or artificial conditions according to various methods well known in the art, and may rely on any known method for the insertion of foreign nucleic acid sequences into a prokarvotic or eukarvotic host cell. The method for transformation is selected

based on the type of host cell being transformed and may include, but is not limited to, viral infection, electroporation, heat shock, lipofection, and particle bombardment. The term "transformed" cells includes stably transformed cells in which the inserted DNA is capable of replication either as an autonomously replicating plasmid or as part of the host chromosome, as well as transiently transformed cells which express the inserted DNA or RNA for limited periods of time.

A "variant" of a particular nucleic acid sequence is defined as a nucleic acid sequence having at least 40% sequence identity to the particular nucleic acid sequence over a certain length of one of the nucleic acid sequences using blastn with the "BLAST 2 Sequences" tool Version 2.0.9 (May-07-1999) set at default parameters. Such a pair of nucleic acids may show, for example, at least 50%, at least 60%, at least 70%, at least 80%, at least 85%, at least 90%, at least 95% or at least 98% or greater sequence identity over a certain defined length. A variant may be described as, for example, an "allelic" (as defined above), "splice," "species," or "polymorphic" variant. A splice variant may have significant identity to a reference molecule, but will generally have a greater or lesser number of polynucleotides due to alternate splicing of exons during mRNA processing. The corresponding polypeptide may possess additional functional domains or lack domains that are present in the reference molecule. Species variants are polynucleotide sequences that vary from one species to another. The resulting polypeptides generally will have significant amino acid identity relative to each other. A polymorphic variant is a variation in the polynucleotide sequence of a particular gene between individuals of a given species. Polymorphic variants also may encompass "single nucleotide polymorphisms" (SNPs) in which the polynucleotide sequence varies by one nucleotide base. The presence of SNPs may be indicative of, for example, a certain population, a disease state, or a propensity for a disease state.

A "variant" of a particular polypeptide sequence is defined as a polypeptide sequence having at least 40% sequence identity to the particular polypeptide sequence over a certain length of one of the polypeptide sequences using blastp with the "BLAST 2 Sequences" tool Version 2.0.9 (May-07-1999) set at default parameters. Such a pair of polypeptides may show, for example, at least 50%, at least 60%, at least 70%, at least 90%, at least 95%, or at least 98% or greater sequence identity over a certain defined length of one of the polypeptides.

30 THE INVENTION

20

35

The invention is based on the discovery of new human nucleic-acid binding proteins (NuABP), the polynucleotides encoding NuABP, and the use of these compositions for the diagnosis, treatment, or prevention of reproductive, immune, and neurological disorders, and cell proliferative disorders including cancer.

Table 1 lists the Incyte clones used to assemble full length nucleotide sequences encoding

NuABP. Columns 1 and 2 show the sequence identification numbers (SEQ ID NOs) of the polypeptide and nucleotide sequences, respectively. Column 3 shows the clone IDs of the Incyte clones in which nucleic acids encoding each NuABP were identified, and column 4 shows the cDNA libraries from which these clones were isolated. Column 5 shows Incyte clones and their corresponding cDNA libraries. Clones for which cDNA libraries are not indicated were derived from pooled cDNA libraries. The Incyte clones in column 5 were used to assemble the consensus nucleotide sequence of each NuABP and are useful as fragments in hybridization technologies.

The columns of Table 2 show various properties of each of the polypeptides of the invention: column 1 references the SEQ ID NO; column 2 shows the number of amino acid residues in each polypeptide; column 3 shows potential phosphorylation sites; column 4 shows potential glycosylation sites; column 5 shows the amino acid residues comprising signature sequences and motifs; column 6 shows identification or homologous sequences as identified by BLAST analysis; and column 7 shows analytical methods and in some cases, searchable databases to which the analytical methods were applied. The methods of column 7 were used to characterize each polypeptide through sequence homology and protein motifs.

The columns of Table 3 show the tissue-specificity and diseases, disorders, or conditions associated with nucleotide sequences encoding NuABP. The first column of Table 3 lists the nucleotide SEQ ID NOs. Column 2 lists fragments of the nucleotide sequences of column 1. These fragments are useful, for example, in hybridization or amplification technologies to identify SEQ ID NO:56-110 and to distinguish between SEQ ID NO:56-110 and related polynucleotide sequences. The polypeptides encoded by these fragments are useful, for example, as immunogenic peptides. Column 3 lists tissue categories which express NuABP as a fraction of total tissues expressing NuABP. Column 4 lists diseases, disorders, or conditions associated with those tissues expressing NuABP as a fraction of total tissues expressing NuABP. Of particular note is the expression of SEQ ID NO:83 and SEQ ID NO:110 in neurological tissue. About 53% of the cDNA libraries expressing SEQ ID NO:83 are derived from neurological tissue. Furthermore, SEQ ID NO:110 expression is detected exclusively in a cDNA library derived from brain tissue afflicted with Huntington's disease. Column 5 lists the vectors used to subclone each cDNA library.

The columns of Table 4 show descriptions of the tissues used to construct the cDNA libraries from which cDNA clones encoding NuABP were isolated. Column 1 references the nucleotide SEQ ID NOs, column 2 shows the cDNA libraries from which these clones were isolated, and column 3 shows the tissue origins and other descriptive information relevant to the cDNA libraries in column 2.

30

Fragments of the nucleotide sequences encoding NuABP are useful, for example, in hybridization or amplification technologies to identify SEQ ID NOS:56-110 and to distinguish

between SEO ID NOS:56-110 and related polynucleotide sequences. The polypeptides encoded by

- Expression line a section of the set for the man most

these fragments are useful, for example, as immunogenic peptides.

20

25

30

35

The invention also encompasses NuABP variants. A preferred NuABP variant is one which has at least about 80%, or alternatively at least about 90%, or even at least about 95% amino acid sequence identity to the NuABP amino acid sequence, and which contains at least one functional or structural characteristic of NuABP.

The invention also encompasses polynucleotides which encode NuABP. In a particular embodiment, the invention encompasses a polynucleotide sequence comprising a sequence selected from the group consisting of SEQ ID NO:56-110, which encodes NuABP.

The invention also encompasses a variant of a polynucleotide sequence encoding NuABP. In particular, such a variant polynucleotide sequence will have at least about 70%, or alternatively at least about 85%, or even at least about 95% polynucleotide sequence identity to the polynucleotide sequence encoding NuABP. A particular aspect of the invention encompasses a variant of a polynucleotide sequence comprising a sequence selected from the group consisting of SEQ ID NO:56-110 which has at least about 70%, or alternatively at least about 85%, or even at least about 95% polynucleotide sequence identity to a nucleic acid sequence selected from the group consisting of SEQ ID NO:56-110. Any one of the polynucleotide variants described above can encode an amino acid sequence which contains at least one functional or structural characteristic of NuABP.

It will be appreciated by those skilled in the art that as a result of the degeneracy of the genetic code, a multitude of polynucleotide sequences encoding NuABP, some bearing minimal similarity to the polynucleotide sequences of any known and naturally occurring gene, may be produced. Thus, the invention contemplates each and every possible variation of polynucleotide sequence that could be made by selecting combinations based on possible codon choices. These combinations are made in accordance with the standard triplet genetic code as applied to the polynucleotide sequence of naturally occurring NuABP, and all such variations are to be considered as being specifically disclosed.

Although nucleotide sequences which encode NuABP and its variants are generally capable of hybridizing to the nucleotide sequence of the naturally occurring NuABP under appropriately selected conditions of stringency, it may be advantageous to produce nucleotide sequences encoding NuABP or its derivatives possessing a substantially different codon usage, e.g., inclusion of non-naturally occurring codons. Codons may be selected to increase the rate at which expression of the peptide occurs in a particular prokaryotic or eukaryotic host in accordance with the frequency with which particular codons are utilized by the host. Other reasons for substantially altering the nucleotide sequence encoding NuABP and its derivatives without altering the encoded amino acid sequences include the production of RNA transcripts having more desirable properties, such as a greater half-life, than transcripts produced from the naturally occurring sequence.

The invention also encompasses production of DNA sequences which encode NuABP and NuABP derivatives, or fragments thereof, entirely by synthetic chemistry. After production, the synthetic sequence may be inserted into any of the many available expression vectors and cell systems using reagents well known in the art. Moreover, synthetic chemistry may be used to introduce mutations into a sequence encoding NuABP or any fragment thereof.

Also encompassed by the invention are polynucleotide sequences that are capable of hybridizing to the claimed polynucleotide sequences, and, in particular, to those shown in SEQ ID NO:56-110 and fragments thereof under various conditions of stringency. (See, e.g., Wahl, G.M. and S.L. Berger (1987) Methods Enzymol. 152:399-407; Kimmel, A.R. (1987) Methods Enzymol. 152:507-511.) Hybridization conditions, including annealing and wash conditions, are described in "Definitions."

Methods for DNA sequencing are well known in the art and may be used to practice any of the embodiments of the invention. The methods may employ such enzymes as the Klenow fragment of DNA polymerase I, SEQUENASE (US Biochemical, Cleveland OH), Taq polymerase (Perkin-Elmer), thermostable T7 polymerase (Amersham Pharmacia Biotech, Piscataway NJ), or combinations of polymerases and proofreading exonucleases such as those found in the ELONGASE amplification system (Life Technologies, Gaithersburg MD). Preferably, sequence preparation is automated with machines such as the MICROLAB 2200 liquid transfer system (Hamilton, Reno NV), PTC200 thermal cycler (MJ Research, Watertown MA) and ABI CATALYST 800 thermal cycler (Perkin-Elmer). Sequencing is then carried out using either the ABI 373 or 377 DNA sequencing system (Perkin-Elmer), the MEGABACE 1000 DNA sequencing system (Molecular Dynamics, Sunnyvale CA), or other systems known in the art. The resulting sequences are analyzed using a variety of algorithms which are well known in the art. (See, e.g., Ausubel, F.M. (1997) Short Protocols in Molecular Biology, John Wiley & Sons, New York NY, unit 7.7; Meyers, R.A. (1995) Molecular Biology and Biotechnology, Wiley VCH, New York NY, pp. 856-853.)

15

20

25

30

The nucleic acid sequences encoding NuABP may be extended utilizing a partial nucleotide sequence and employing various PCR-based methods known in the art to detect upstream sequences, such as promoters and regulatory elements. For example, one method which may be employed, restriction-site PCR, uses universal and nested primers to amplify unknown sequence from genomic DNA within a cloning vector. (See, e.g., Sarkar, G. (1993) PCR Methods Applic. 2:318-322.)

Another method, inverse PCR, uses primers that extend in divergent directions to amplify unknown sequence from a circularized template. The template is derived from restriction fragments comprising a known genomic locus and surrounding sequences. (See, e.g., Triglia, T. et al. (1988) Nucleic Acids Res. 16:8186.) A third method, capture PCR, involves PCR amplification of DNA fragments adjacent to known sequences in human and yeast artificial chromosome DNA. (See, e.g., Lagerstrom, M. et al.

(1991) PCR Methods Applic. 1:111-119.) In this method, multiple restriction enzyme digestions and ligations may be used to insert an engineered double-stranded sequence into a region of unknown sequence before performing PCR. Other methods which may be used to retrieve unknown sequences are known in the art. (See, e.g., Parker, J.D. et al. (1991) Nucleic Acids Res. 19:3055-3060). Additionally, one may use PCR, nested primers, and PROMOTERFINDER libraries (Clontech, Palo Alto CA) to walk genomic DNA. This procedure avoids the need to screen libraries and is useful in finding intron/exon junctions. For all PCR-based methods, primers may be designed using commercially available software, such as OLIGO 4.06 Primer Analysis software (National Biosciences, Plymouth MN) or another appropriate program, to be about 22 to 30 nucleotides in length, to have a GC content of about 50% or more, and to anneal to the template at temperatures of about 68°C to 72°C.

When screening for full-length cDNAs, it is preferable to use libraries that have been size-selected to include larger cDNAs. In addition, random-primed libraries, which often include sequences containing the 5' regions of genes, are preferable for situations in which an oligo d(T) library does not yield a full-length cDNA. Genomic libraries may be useful for extension of sequence into 5' non-transcribed regulatory regions.

10

15

20

30

Capillary electrophoresis systems which are commercially available may be used to analyze the size or confirm the nucleotide sequence of sequencing or PCR products. In particular, capillary sequencing may employ flowable polymers for electrophoretic separation, four different nucleotide-specific, laser-stimulated fluorescent dyes, and a charge coupled device camera for detection of the emitted wavelengths. Output/light intensity may be converted to electrical signal using appropriate software (e.g., GENOTYPER and SEQUENCE NAVIGATOR, Perkin-Elmer), and the entire process from loading of samples to computer analysis and electronic data display may be computer controlled. Capillary electrophoresis is especially preferable for sequencing small DNA fragments which may be present in limited amounts in a particular sample.

In another embodiment of the invention, polynucleotide sequences or fragments thereof which encode NuABP may be cloned in recombinant DNA molecules that direct expression of NuABP, or fragments or functional equivalents thereof, in appropriate host cells. Due to the inherent degeneracy of the genetic code, other DNA sequences which encode substantially the same or a functionally equivalent amino acid sequence may be produced and used to express NuABP.

The nucleotide sequences of the present invention can be engineered using methods generally known in the art in order to alter NuABP-encoding sequences for a variety of purposes including, but not limited to, modification of the cloning, processing, and/or expression of the gene product. DNA shuffling by random fragmentation and PCR reassembly of gene fragments and synthetic

35 oligonucleotides may be used to engineer the nucleotide sequences. For example, oligonucleotide-

mediated site-directed mutagenesis may be used to introduce mutations that create new restriction sites, alter glycosylation patterns, change codon preference, produce splice variants, and so forth.

In another embodiment, sequences encoding NuABP may be synthesized, in whole or in part, using chemical methods well known in the art. (See, e.g., Caruthers, M.H. et al. (1980) Nucleic Acids Symp. Ser. 7:215-223; and Horn, T. et al. (1980) Nucleic Acids Symp. Ser. 7:225-232.)

Alternatively, NuABP itself or a fragment thereof may be synthesized using chemical methods. For example, peptide synthesis can be performed using various solid-phase techniques. (See, e.g., Roberge, J.Y. et al. (1995) Science 269:202-204.) Automated synthesis may be achieved using the ABI 431A peptide synthesizer (Perkin-Elmer). Additionally, the amino acid sequence of NuABP, or any part thereof, may be altered during direct synthesis and/or combined with sequences from other proteins, or any part thereof, to produce a variant polypeptide.

The peptide may be substantially purified by preparative high performance liquid chromatography. (See, e.g., Chiez, R.M. and F.Z. Regnier (1990) Methods Enzymol. 182:392-421.)

The composition of the synthetic peptides may be confirmed by amino acid analysis or by sequencing. (See, e.g., Creighton, T. (1984) Proteins, Structures and Molecular Properties. WH Freeman, New York NY.)

In order to express a biologically active NuABP, the nucleotide sequences encoding NuABP or derivatives thereof may be inserted into an appropriate expression vector, i.e., a vector which contains the necessary elements for transcriptional and translational control of the inserted coding sequence in a suitable host. These elements include regulatory sequences, such as enhancers, constitutive and inducible promoters, and 5' and 3' untranslated regions in the vector and in polynucleotide sequences encoding NuABP. Such elements may vary in their strength and specificity. Specific initiation signals may also be used to achieve more efficient translation of sequences encoding NuABP. Such signals include the ATG initiation codon and adjacent sequences. e.g. the Kozak sequence. In cases where sequences encoding NuABP and its initiation codon and upstream regulatory sequences are inserted into the appropriate expression vector, no additional transcriptional or translational control signals may be needed. However, in cases where only coding sequence, or a fragment thereof, is inserted, exogenous translational control signals including an inframe ATG initiation codon should be provided by the vector. Exogenous translational elements and initiation codons may be of various origins, both natural and synthetic. The efficiency of expression may be enhanced by the inclusion of enhancers appropriate for the particular host cell system used. (See, e.g., Scharf, D. et al. (1994) Results Probl. Cell Differ. 20:125-162.)

20

30

35

Methods which are well known to those skilled in the art may be used to construct expression vectors containing sequences encoding NuABP and appropriate transcriptional and translational control elements. These methods include in vitro recombinant DNA techniques, synthetic techniques,

WO 00/44900

5

10

20

25

30

PCT/US00/02237

and in vivo genetic recombination. (See, e.g., Sambrook, J. et al. (1989) Molecular Cloning, A
Laboratory Manual, Cold Spring Harbor Press, Plainview NY, ch. 4, 8, and 16-17; Ausubel, F.M. et
al. (1995) Current Protocols in Molecular Biology. John Wiley & Sons, New York NY, ch. 9, 13, and
16.)

A variety of expression vector/host systems may be utilized to contain and express sequences encoding NuABP. These include, but are not limited to, microorganisms such as bacteria transformed with recombinant bacteriophage, plasmid, or cosmid DNA expression vectors; yeast transformed with yeast expression vectors; insect cell systems infected with viral expression vectors (e.g., baculovirus); plant cell systems transformed with viral expression vectors (e.g., cauliflower mosaic virus, CaMV, or tobacco mosaic virus, TMV) or with bacterial expression vectors (e.g., Ti or pBR322 plasmids); or animal cell systems. The invention is not limited by the host cell employed.

In bacterial systems, a number of cloning and expression vectors may be selected depending upon the use intended for polynucleotide sequences encoding NuABP. For example, routine cloning, subcloning, and propagation of polynucleotide sequences encoding NuABP can be achieved using a multifunctional E. coli vector such as PBLUESCRIPT (Stratagene, La Jolla CA) or PSPORT1 plasmid (Life Technologies). Ligation of sequences encoding NuABP into the vector's multiple cloning site disrupts the *lacZ* gene, allowing a colorimetric screening procedure for identification of transformed bacteria containing recombinant molecules. In addition, these vectors may be useful for in vitro transcription, dideoxy sequencing, single strand rescue with helper phage, and creation of nested deletions in the cloned sequence. (See, e.g., Van Heeke, G. and S.M. Schuster (1989) J. Biol. Chem. 264:5503-5509.) When large quantities of NuABP are needed, e.g. for the production of antibodies, vectors which direct high level expression of NuABP may be used. For example, vectors containing the strong, inducible T5 or T7 bacteriophage promoter may be used.

Yeast expression systems may be used for production of NuABP. A number of vectors containing constitutive or inducible promoters, such as alpha factor, alcohol oxidase, and PGH promoters, may be used in the yeast Saccharomyces cerevisiae or Pichia pastoris. In addition, such vectors direct either the secretion or intracellular retention of expressed proteins and enable integration of foreign sequences into the host genome for stable propagation. (See, e.g., Ausubel, 1995, supra; Bitter, G.A. et al. (1987) Methods Enzymol. 153:516-544; and Scorer, C.A. et al. (1994) Bio/Technology 12:181-184.)

Plant systems may also be used for expression of NuABP. Transcription of sequences encoding NuABP may be driven viral promoters, e.g., the 35S and 19S promoters of CaMV used alone or in combination with the omega leader sequence from TMV (Takamatsu, N. (1987) EMBO J. 6:307-311). Alternatively, plant promoters such as the small subunit of RUBISCO or heat shock promoters may be used. (See, e.g., Coruzzi, G. et al. (1984) EMBO J. 3:1671-1680; Broglie, R. et al.

(1984) Science 224:838-843; and Winter, J. et al. (1991) Results Probl. Cell Differ. 17:85-105.)
These constructs can be introduced into plant cells by direct DNA transformation or pathogen-mediated transfection. (See, e.g., <u>The McGraw Hill Yearbook of Science and Technology</u> (1992) McGraw Hill, New York NY, pp. 191-196.)

5

25

In mammalian cells, a number of viral-based expression systems may be utilized. In cases where an adenovirus is used as an expression vector, sequences encoding NuABP may be ligated into an adenovirus transcription/translation complex consisting of the late promoter and tripartite leader sequence. Insertion in a non-essential E1 or E3 region of the viral genome may be used to obtain infective virus which expresses NuABP in host cells. (See, e.g., Logan, J. and T. Shenk (1984) Proc. Natl. Acad. Sci. USA 81:3655-3659.) In addition, transcription enhancers, such as the Rous sarcoma virus (RSV) enhancer, may be used to increase expression in mammalian host cells. SV40 or EBV-based vectors may also be used for high-level protein expression.

Human artificial chromosomes (HACs) may also be employed to deliver larger fragments of DNA than can be contained in and expressed from a plasmid. HACs of about 6 kb to 10 Mb are constructed and delivered via conventional delivery methods (liposomes, polycationic amino polymers, or vesicles) for therapeutic purposes. (See, e.g., Harrington, J.J. et al. (1997) Nat. Genet. 15:345-355.)

For long term production of recombinant proteins in mammalian systems, stable expression of NuABP in cell lines is preferred. For example, sequences encoding NuABP can be transformed into cell lines using expression vectors which may contain viral origins of replication and/or endogenous expression elements and a selectable marker gene on the same or on a separate vector. Following the introduction of the vector, cells may be allowed to grow for about 1 to 2 days in enriched media before being switched to selective media. The purpose of the selectable marker is to confer resistance to a selective agent, and its presence allows growth and recovery of cells which successfully express the introduced sequences. Resistant clones of stably transformed cells may be propagated using tissue culture techniques appropriate to the cell type.

Any number of selection systems may be used to recover transformed cell lines. These include, but are not limited to, the herpes simplex virus thymidine kinase and adenine phosphoribosyltransferase genes, for use in the and apr cells, respectively. (See, e.g., Wigler, M. et al. (1977) Cell 11:223-232; Lowy, I. et al. (1980) Cell 22:817-823.) Also, antimetabolite, antibiotic, or herbicide resistance can be used as the basis for selection. For example, dhfr confers resistance to methotrexate; neo confers resistance to the aminoglycosides neomycin and G-418; and als and pat confer resistance to chlorsulfuron and phosphinotricin acetyltransferase, respectively. (See, e.g., Wigler, M. et al. (1980) Proc. Natl. Acad. Sci. USA 77:3567-3570; Colbere-Garapin, F. et al. (1981) J. Mol. Biol. 150:1-14.) Additional selectable genes have been described, e.g., trpB and hisD, which

alter cellular requirements for metabolites. (See, e.g., Hartman, S.C. and R.C. Mulligan (1988) Proc. Natl. Acad. Sci. USA 85:8047-8051.) Visible markers, e.g., anthocyanins, green fluorescent proteins (GFP: Clontech), β glucuronidase and its substrate β-glucuronide, or luciferase and its substrate luciferin may be used. These markers can be used not only to identify transformants, but also to quantify the amount of transient or stable protein expression attributable to a specific vector system. (See, e.g., Rhodes, C.A. (1995) Methods Mol. Biol. 55:121-131.)

Although the presence/absence of marker gene expression suggests that the gene of interest is also present, the presence and expression of the gene may need to be confirmed. For example, if the sequence encoding NuABP is inserted within a marker gene sequence, transformed cells containing sequences encoding NuABP can be identified by the absence of marker gene function. Alternatively, a marker gene can be placed in tandem with a sequence encoding NuABP under the control of a single promoter. Expression of the marker gene in response to induction or selection usually indicates expression of the tandem gene as well.

10

15

20

30

35

In general, host cells that contain the nucleic acid sequence encoding NuABP and that express NuABP may be identified by a variety of procedures known to those of skill in the art. These procedures include, but are not limited to, DNA-DNA or DNA-RNA hybridizations, PCR amplification, and protein bioassay or immunoassay techniques which include membrane, solution, or chip based technologies for the detection and/or quantification of nucleic acid or protein sequences.

Immunological methods for detecting and measuring the expression of NuABP using either specific polyclonal or monoclonal antibodies are known in the art. Examples of such techniques include enzyme-linked immunosorbent assays (ELISAs), radioimmunoassays (RIAs), and fluorescence activated cell sorting (FACS). A two-site, monoclonal-based immunoassay utilizing monoclonal antibodies reactive to two non-interfering epitopes on NuABP is preferred, but a competitive binding assay may be employed. These and other assays are well known in the art. (See, e.g., Hampton, R. et al. (1990) Serological Methods, a Laboratory Manual, APS Press, St. Paul MN, 25 Sect. IV; Coligan, J.E. et al. (1997) Current Protocols in Immunology, Greene Pub. Associates and Wiley-Interscience, New York NY; and Pound, J.D. (1998) Immunochemical Protocols, Humana Press, Totowa NJ.)

A wide variety of labels and conjugation techniques are known by those skilled in the art and may be used in various nucleic acid and amino acid assays. Means for producing labeled hybridization or PCR probes for detecting sequences related to polynucleotides encoding NuABP include oligolabeling, nick translation, end-labeling, or PCR amplification using a labeled nucleotide. Alternatively, the sequences encoding NuABP, or any fragments thereof, may be cloned into a vector for the production of an mRNA probe. Such vectors are known in the art, are commercially available. and may be used to synthesize RNA probes in vitro by addition of an appropriate RNA polymerase

such as T7. T3, or SP6 and labeled nucleotides. These procedures may be conducted using a variety of commercially available kits, such as those provided by Amersham Pharmacia Biotech. Promega (Madison WI), and US Biochemical. Suitable reporter molecules or labels which may be used for ease of detection include radionuclides, enzymes, fluorescent, chemiluminescent, or chromogenic agents, as well as substrates, cofactors, inhibitors, magnetic particles, and the like.

Host cells transformed with nucleotide sequences encoding NuABP may be cultured under conditions suitable for the expression and recovery of the protein from cell culture. The protein produced by a transformed cell may be secreted or retained intracellularly depending on the sequence and/or the vector used. As will be understood by those of skill in the art, expression vectors containing polynucleotides which encode NuABP may be designed to contain signal sequences which direct secretion of NuABP through a prokaryotic or cukaryotic cell membrane.

In addition, a host cell strain may be chosen for its ability to modulate expression of the inserted sequences or to process the expressed protein in the desired fashion. Such modifications of the polypeptide include, but are not limited to, acetylation, carboxylation, glycosylation, phosphorylation, lipidation, and acylation. Post-translational processing which cleaves a "prepro" or "pro" form of the protein may also be used to specify protein targeting, folding, and/or activity. Different host cells which have specific cellular machinery and characteristic mechanisms for post-translational activities (e.g., CHO, HeLa, MDCK, HEK293, and W138) are available from the American Type Culture Collection (ATCC, Manassas VA) and may be chosen to ensure the correct modification and processing of the foreign protein.

20

35

In another embodiment of the invention. natural, modified, or recombinant nucleic acid sequences encoding NuABP may be ligated to a heterologous sequence resulting in translation of a fusion protein in any of the aforementioned host systems. For example, a chimeric NuABP protein containing a heterologous moiety that can be recognized by a commercially available antibody may facilitate the screening of peptide libraries for inhibitors of NuABP activity. Heterologous protein and peptide moieties may also facilitate purification of fusion proteins using commercially available affinity matrices. Such moieties include, but are not limited to, glutathione S-transferase (GST), maltose binding protein (MBP), thioredoxin (Trx), calmodulin binding peptide (CBP), 6-His, FLAG, c-myc, and hemagglutinin (HA). GST, MBP, Trx, CBP, and 6-His enable purification of their cognate fusion proteins on immobilized glutathione, maltose, phenylarsine oxide, calmodulin, and metal-chelate resins, respectively. FLAG, c-myc, and hemagglutinin (HA) enable immunoaffinity purification of fusion proteins using commercially available monoclonal and polyclonal antibodies that specifically recognize these epitope tags. A fusion protein may also be engineered to contain a proteolytic cleavage site located between the NuABP encoding sequence and the heterologous protein sequence, so that NuABP may be cleaved away from the heterologous moiety following purification.

Methods for fusion protein expression and purification are discussed in Ausubel (1995, supra, ch. 10).

A variety of commercially available kits may also be used to facilitate expression and purification of fusion proteins.

In a further embodiment of the invention, synthesis of radiolabeled NuABP may be achieved in vitro using the TNT rabbit reticulocyte lysate or wheat germ extract system (Promega). These systems couple transcription and translation of protein-coding sequences operably associated with the T7, T3, or SP6 promoters. Translation takes place in the presence of a radiolabeled amino acid precursor, for example, 35S-methionine.

Fragments of NuABP may be produced not only by recombinant means, but also by direct peptide synthesis using solid-phase techniques. (See, e.g., Creighton, supra, pp. 55-60.) Protein synthesis may be performed by manual techniques or by automation. Automated synthesis may be achieved, for example, using the ABI 431A peptide synthesizer (Perkin-Elmer). Various fragments of NuABP may be synthesized separately and then combined to produce the full length molecule.

THERAPEUTICS

10

15

20

Chemical and structural similarity, e.g., in the context of sequences and motifs, exists between regions of NuABP and nucleic-acid binding proteins. In addition, the expression of NuABP is closely associated with proliferative, neuronal, inflamed, and cancerous tissues and tissues of the reproductive system. Therefore, NuABP appears to play a role in reproductive, immune, and neurological disorders, and cell proliferative disorders including cancer. In the treatment of disorders associated with increased NuABP expression or activity, it is desirable to decrease the expression or activity of NuABP. In the treatment of disorders associated with decreased NuABP expression or activity, it is desirable to increase the expression or activity of NuABP.

Therefore, in one embodiment, NuABP or a fragment or derivative thereof may be administered to a subject to treat or prevent a disorder associated with decreased expression or activity of NuABP. Examples of such disorders include, but are not limited to, a reproductive disorder such as disorders of prolactin production, infertility, including tubal disease, ovulatory defects, and endometriosis, disruptions of the estrous cycle, disruptions of the menstrual cycle, polycystic ovary syndrome, ovarian hyperstimulation syndrome, endometrial and ovarian tumors, uterine fibroids, autoimmune disorders, ectopic pregnancies, and teratogenesis; cancer of the breast, fibrocystic breast disease, and galactorrhea; disruptions of spermatogenesis, abnormal sperm physiology, cancer of the testis, cancer of the prostate, benign prostatic hyperplasia, prostatitis, Peyronie's disease, impotence, carcinoma of the male breast, and gynecomastia; an immune disorder such as inflammation, actinic keratosis, acquired immunodeficiency syndrome (AIDS), Addison's disease, adult respiratory distress syndrome, allergies, ankylosing spondylitis, amyloidosis, anemia, arteriosclerosis. asthma,

cholecystitis, cirrhosis, contact dermatitis, Crohn's disease, atopic dermatitis, dermatomyositis, diabetes mellitus, emphysema, erythroblastosis fetalis, erythema nodosum, atrophic gastritis, glomerulonephritis. Goodpasture's syndrome, gout, Graves' disease. Hashimoto's thyroiditis, paroxysmal nocturnal hemoglobinuria, hepatitis, hypereosinophilia, irritable bowel syndrome, episodic lymphopenia with lymphocytotoxins, mixed connective tissue disease (MCTD), multiple sclerosis, myasthenia gravis, myocardial or pericardial inflammation, myelofibrosis, osteoarthritis, osteoporosis, pancreatitis, polycythemia vera, polymyositis, psoriasis, Reiter's syndrome, rheumatoid arthritis, scleroderma. Sjögren's syndrome, systemic anaphylaxis, systemic lupus erythematosus, systemic sclerosis, primary thrombocythemia, thrombocytopenic purpura, ulcerative colitis, uveitis, Werner syndrome, complications of cancer, hemodialysis, and extracorporeal circulation, trauma, and hematopoietic cancer including lymphoma, leukemia, and myeloma; a neurological disorder such as epilepsy, ischemic cerebrovascular disease, stroke, cerebral neoplasms, Alzheimer's disease, Pick's disease, Huntington's disease, dementia, Parkinson's disease and other extrapyramidal disorders, amyotrophic lateral sclerosis and other motor neuron disorders, progressive neural muscular atrophy, retinitis pigmentosa, hereditary ataxias, multiple sclerosis and other demyelinating diseases, bacterial and viral meningitis, brain abscess, subdural empyema, epidural abscess, suppurative intracranial thrombophlebitis, myelitis and radiculitis, viral central nervous system disease, prion diseases including kuru, Creutzfeldt-Jakob disease, and Gerstmann-Straussler-Scheinker syndrome, fatal familial insomnia, nutritional and metabolic diseases of the nervous system, neurofibromatosis, tuberous sclerosis, cerebelloretinal hemangioblastomatosis, encephalotrigeminal syndrome, mental retardation and other developmental disorders of the central nervous system, cerebral palsy, neuroskeletal disorders, autonomic nervous system disorders, cranial nerve disorders, spinal cord diseases, muscular dystrophy and other neuromuscular disorders, peripheral nervous system disorders, dermatomyositis and polymyositis, inherited, metabolic, endocrine, and toxic myopathies, myasthenia gravis, periodic paralysis, mental disorders including mood, anxiety, and schizophrenic disorders, akathesia, amnesia, catatonia, diabetic neuropathy, tardive dyskinesia, dystonias, paranoid psychoses, postherpetic neuralgia, and Tourette's disorder; and a cell proliferative disorder such as actinic keratosis, arteriosclerosis, atherosclerosis, bursitis, cirrhosis, hepatitis, mixed connective tissue disease (MCTD), myelofibrosis, paroxysmal nocturnal hemoglobinuria, polycythemia vera, psoriasis, primary thrombocythemia, and cancers including adenocarcinoma, leukemia, lymphoma, melanoma, myeloma, sarcoma, teratocarcinoma, and, in particular, cancers of the adrenal gland, bladder, bone, bone marrow, brain, breast, cervix, gall bladder, ganglia, gastrointestinal tract, heart, kidney, liver, lung, muscle, ovary, pancreas, parathyroid, penis, prostate, salivary glands, skin, spleen, testis, thymus, thyroid, and uterus.

10

20

35

In another embodiment, a vector capable of expressing NuABP or a fragment or derivative

thereof may be administered to a subject to treat or prevent a disorder associated with decreased expression or activity of NnABP including, but not limited to, those described above.

In a further embodiment, a pharmaceutical composition comprising a substantially purified NuABP in conjunction with a suitable pharmaceutical carrier may be administered to a subject to treat or prevent a disorder associated with decreased expression or activity of NuABP including, but not limited to, those provided above.

In still another embodiment, an agonist which modulates the activity of NuABP may be administered to a subject to treat or prevent a disorder associated with decreased expression or activity of NuABP including, but not limited to, those listed above.

10

15

20

25

35

In a further embodiment, an antagonist of NuABP may be administered to a subject to treat or prevent a disorder associated with increased expression or activity of NuABP. Examples of such disorders include, but are not limited to, those reproductive, immune, and neurological disorders, and cell proliferative disorders including cancer, described above. In one aspect, an antibody which specifically binds NuABP may be used directly as an antagonist or indirectly as a targeting or delivery mechanism for bringing a pharmaceutical agent to cells or tissues which express NuABP.

In an additional embodiment, a vector expressing the complement of the polynucleotide encoding NuABP may be administered to a subject to treat or prevent a disorder associated with increased expression or activity of NuABP including, but not limited to, those described above.

In other embodiments, any of the proteins, antagonists. antibodies, agonists, complementary sequences, or vectors of the invention may be administered in combination with other appropriate therapeutic agents. Selection of the appropriate agents for use in combination therapy may be made by one of ordinary skill in the art, according to conventional pharmaceutical principles. The combination of therapeutic agents may act synergistically to effect the treatment or prevention of the various disorders described above. Using this approach, one may be able to achieve therapeutic efficacy with lower dosages of each agent, thus reducing the potential for adverse side effects.

An antagonist of NuABP may be produced using methods which are generally known in the art. In particular, purified NuABP may be used to produce antibodies or to screen libraries of pharmaceutical agents to identify those which specifically bind NuABP. Antibodies to NuABP may also be generated using methods that are well known in the art. Such antibodies may include, but are not limited to, polyclonal, monoclonal, chimeric, and single chain antibodies, Fab fragments, and fragments produced by a Fab expression library. Neutralizing antibodies (i.e., those which inhibit dimer formation) are generally preferred for therapeutic use.

For the production of antibodies, various hosts including goats, rabbits, rats, mice, humans, and others may be immunized by injection with NuABP or with any fragment or oligopeptide thereof which has immunogenic properties. Depending on the host species, various adjuvants may be used to

increase immunological response. Such adjuvants include, but are not limited to, Freund's, mineral gets such as aluminum hydroxide, and surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, KLH, and dinitrophenol. Among adjuvants used in humans. BCG (bacilli Calmette-Guerin) and Corynebacterium parvum are especially preferable.

It is preferred that the oligopeptides, peptides, or fragments used to induce antibodies to NuABP have an amino acid sequence consisting of at least about 5 amino acids, and generally will consist of at least about 10 amino acids. It is also preferable that these oligopeptides, peptides, or fragments are identical to a portion of the amino acid sequence of the natural protein and contain the entire amino acid sequence of a small, naturally occurring molecule. Short stretches of NuABP amino acids may be fused with those of another protein, such as KLH, and antibodies to the chimeric molecule may be produced.

Monoclonal antibodies to NuABP may be prepared using any technique which provides for the production of antibody molecules by continuous cell lines in culture. These include, but are not limited to, the hybridoma technique, the human B-cell hybridoma technique, and the EBV-hybridoma technique. (See, e.g., Kohler, G. et al. (1975) Nature 256:495-497: Kozbor, D. et al. (1985) J. Immunol. Methods 81:31-42; Cote, R.J. et al. (1983) Proc. Natl. Acad. Sci. USA 80:2026-2030; and Cole, S.P. et al. (1984) Mol. Cell Biol. 62:109-120.)

In addition, techniques developed for the production of "chimeric antibodies," such as the splicing of mouse antibody genes to human antibody genes to obtain a molecule with appropriate antigen specificity and biological activity, can be used. (See, e.g., Morrison, S.L. et al. (1984) Proc. Natl. Acad. Sci. USA 81:6851-6855; Neuberger, M.S. et al. (1984) Nature 312:604-608; and Takeda, S. et al. (1985) Nature 314:452-454.) Alternatively, techniques described for the production of single chain antibodies may be adapted, using methods known in the art, to produce NuABP-specific single chain antibodies. Antibodies with related specificity, but of distinct idiotypic composition, may be generated by chain shuffling from random combinatorial immunoglobulin libraries. (See, e.g., Burton, D.R. (1991) Proc. Natl. Acad. Sci. USA 88:10134-10137.)

Antibodies may also be produced by inducing <u>in vivo</u> production in the lymphocyte population or by screening immunoglobulin libraries or panels of highly specific binding reagents as disclosed in the literature. (See, e.g., Orlandi, R. et al. (1989) Proc. Natl. Acad. Sci. USA 86:3833-3837; Winter, G. et al. (1991) Nature 349:293-299.)

30

35

Antibody fragments which contain specific binding sites for NuABP may also be generated. For example, such fragments include, but are not limited to, F(ab')₂ fragments produced by pepsin digestion of the antibody molecule and Fab fragments generated by reducing the disulfide bridges of the F(ab')₂ fragments. Alternatively, Fab expression libraries may be constructed to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity. (See, e.g., Huse, W.D.

WO 00/44900

10

15

20

25

30

35

PCT/US00/02237

et al. (1989) Science 246:1275-1281.)

Various immunoassays may be used for screening to identify antibodies having the desired specificity. Numerous protocols for competitive binding or immunoradiometric assays using either polyclonal or monoclonal antibodies with established specificities are well known in the art. Such immunoassays typically involve the measurement of complex formation between NuABP and its specific antibody. A two-site, monoclonal-based immunoassay utilizing monoclonal antibodies reactive to two non-interfering NuABP epitopes is generally used, but a competitive binding assay may also be employed (Pound, supra).

Various methods such as Scatchard analysis in conjunction with radioimmunoassay techniques may be used to assess the affinity of antibodies for NuABP. Affinity is expressed as an association constant, K_s, which is defined as the molar concentration of NuABP-antibody complex divided by the molar concentrations of free antigen and free antibody under equilibrium conditions. The K_s determined for a preparation of polyclonal antibodies, which are heterogeneous in their affinities for multiple NuABP epitopes, represents the average affinity, or avidity, of the antibodies for NuABP. The K_s determined for a preparation of monoclonal antibodies, which are monospecific for a particular NuABP epitope, represents a true measure of affinity. High-affinity antibody preparations with K_s ranging from about 10° to 10¹2 L/mole are preferred for use in immunoassays in which the NuABP-antibody complex must withstand rigorous manipulations. Low-affinity antibody preparations with K_s ranging from about 10° to 10² L/mole are preferred for use in immunopurification and similar procedures which ultimately require dissociation of NuABP, preferably in active form, from the antibody (Catty, D. (1988) Antibodies, Volume 1: A Practical Approach, IRL Press, Washington, DC; Liddell, J.E. and Cryer, A. (1991) A Practical Guide to Monoclonal Antibodies, John Wiley & Sons, New York NY).

The titer and avidity of polyclonal antibody preparations may be further evaluated to determine the quality and suitability of such preparations for certain downstream applications. For example, a polyclonal antibody preparation containing at least 1-2 mg specific antibody/ml, preferably 5-10 mg specific antibody/ml, is generally employed in procedures requiring precipitation of NuABP-antibody complexes. Procedures for evaluating antibody specificity, titer, and avidity, and guidelines for antibody quality and usage in various applications, are generally available. (See, e.g., Catty, supra, and Coligan et al. supra.)

In another embodiment of the invention, the polynucleotides encoding NuABP, or any fragment or complement thereof, may be used for therapeutic purposes. In one aspect, the complement of the polynucleotide encoding NuABP may be used in situations in which it would be desirable to block the transcription of the mRNA. In particular, cells may be transformed with sequences complementary to polynucleotides encoding NuABP. Thus, complementary molecules or

fragments may be used to modulate NuABP activity, or to achieve regulation of gene function. Such technology is now well known in the art, and sense or antisense oligonucleotides or larger fragments can be designed from various locations along the coding or control regions of sequences encoding NuABP.

5

10

15

20

30

Expression vectors derived from retroviruses, adenoviruses, or herpes or vaccinia viruses, or from various bacterial plasmids, may be used for delivery of nucleotide sequences to the targeted organ, tissue, or cell population. Methods which are well known to those skilled in the art can be used to construct vectors to express nucleic acid sequences complementary to the polynucleotides encoding NuABP. (See, e.g., Sambrook, supra; Ausubel, 1995, supra.)

Genes encoding NuABP can be turned off by transforming a cell or tissue with expression vectors which express high levels of a polynucleotide, or fragment thereof, encoding NuABP. Such constructs may be used to introduce untranslatable sense or antisense sequences into a cell. Even in the absence of integration into the DNA, such vectors may continue to transcribe RNA molecules until they are disabled by endogenous nucleases. Transient expression may last for a month or more with a non-replicating vector, and may last even longer if appropriate replication elements are part of the vector system.

As mentioned above, modifications of gene expression can be obtained by designing complementary sequences or antisense molecules (DNA, RNA, or PNA) to the control, 5', or regulatory regions of the gene encoding NuABP. Oligonucleotides derived from the transcription initiation site, e.g., between about positions -10 and +10 from the start site, may be employed. Similarly, inhibition can be achieved using triple helix base-pairing methodology. Triple helix pairing is useful because it causes inhibition of the ability of the double helix to open sufficiently for the binding of polymerases. transcription factors, or regulatory molecules. Recent therapeutic advances using triplex DNA have been described in the literature. (See, e.g., Gee, J.E. et al. (1994) in Huber, B.E. and B.I. Carr, Molecular and Immunologic Approaches, Futura Publishing, Mt. Kisco NY, pp. 163-177.) A complementary sequence or antisense molecule may also be designed to block translation of mRNA by preventing the transcript from binding to ribosomes.

Ribozymes, enzymatic RNA molecules, may also be used to catalyze the specific cleavage of RNA. The mechanism of ribozyme action involves sequence-specific hybridization of the ribozyme molecule to complementary target RNA, followed by endonucleolytic cleavage. For example, engineered hammerhead motif ribozyme molecules may specifically and efficiently catalyze endonucleolytic cleavage of sequences encoding NuABP.

Specific ribozyme cleavage sites within any potential RNA target are initially identified by scanning the target molecule for ribozyme cleavage sites, including the following sequences: GUA, GUU, and GUC. Once identified, short RNA sequences of between 15 and 20 ribonucleotides,

corresponding to the region of the target gene containing the cleavage site, may be evaluated for secondary structural features which may render the oligonucleotide inoperable. The suitability of candidate targets may also be evaluated by testing accessibility to hybridization with complementary oligonucleotides using ribonuclease protection assays.

Complementary ribonucleic acid molecules and ribozymes of the invention may be prepared by any method known in the art for the synthesis of nucleic acid molecules. These include techniques for chemically synthesizing oligonucleotides such as solid phase phosphoramidite chemical synthesis. Alternatively, RNA molecules may be generated by in vitro and in vivo transcription of DNA sequences encoding NuABP. Such DNA sequences may be incorporated into a wide variety of vectors with suitable RNA polymerase promoters such as T7 or SP6. Alternatively, these cDNA constructs that synthesize complementary RNA, constitutively or inducibly, can be introduced into cell lines, cells, or tissues.

RNA molecules may be modified to increase intracellular stability and half-life. Possible modifications include, but are not limited to, the addition of flanking sequences at the 5' and/or 3' ends of the molecule, or the use of phosphorothioate or 2' O-methyl rather than phosphodiesterase linkages within the backbone of the molecule. This concept is inherent in the production of PNAs and can be extended in all of these molecules by the inclusion of nontraditional bases such as inosine, queosine, and wybutosine, as well as acetyl-, methyl-, thio-, and similarly modified forms of adenine, cytidine, guanine, thymine, and uridine which are not as easily recognized by endogenous endonucleases.

15

20

25

35

Many methods for introducing vectors into cells or tissues are available and equally suitable for use in vivo, in vitro, and ex vivo. For ex vivo therapy, vectors may be introduced into stem cells taken from the patient and clonally propagated for autologous transplant back into that same patient. Delivery by transfection, by liposome injections, or by polycationic amino polymers may be achieved using methods which are well known in the art. (See, e.g., Goldman, C.K. et al. (1997) Nat. Biotechnol. 15:462-466.)

Any of the therapeutic methods described above may be applied to any subject in need of such therapy, including, for example, mammals such as humans, dogs, cats, cows, horses, rabbits, and monkeys.

An additional embodiment of the invention relates to the administration of a pharmaceutical or sterile composition, in conjunction with a pharmaceutically acceptable carrier, for any of the therapeutic effects discussed above. Such pharmaceutical compositions may consist of NuABP, antibodies to NuABP, and mimetics, agonists, antagonists, or inhibitors of NuABP. The compositions may be administered alone or in combination with at least one other agent, such as a stabilizing compound, which may be administered in any sterile, biocompatible pharmaceutical carrier including, but not limited to, saline, buffered saline, dextrose, and water. The compositions may be administered

to a patient alone, or in combination with other agents, drugs, or hormones.

10

15

20

25

30

35

The pharmaceutical compositions utilized in this invention may be administered by any number of routes including, but not limited to, oral, intravenous, intramuscular, intra-arterial, intramedullary, intrathecal, intraventricular, transdermal, subcutaneous, intraperitoneal, intranasal, enteral, topical, sublingual or rectal means.

In addition to the active ingredients, these pharmaceutical compositions may contain suitable pharmaceutically-acceptable carriers comprising excipients and auxiliaries which facilitate processing of the active compounds into preparations which can be used pharmaceutically. Further details on techniques for formulation and administration may be found in the latest edition of Remington's Pharmaceutical Sciences (Maack Publishing, Easton PA).

Pharmaceutical compositions for oral administration can be formulated using pharmaceutically acceptable carriers well known in the art in dosages suitable for oral administration. Such carriers enable the pharmaceutical compositions to be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions, and the like, for ingestion by the patient.

Pharmaceutical preparations for oral use can be obtained through combining active compounds with solid excipient and processing the resultant mixture of granules (optionally, after grinding) to obtain tablets or dragee cores. Suitable auxiliaries can be added, if desired. Suitable excipients include carbohydrate or protein fillers, such as sugars, including lactose, sucrose, mannitol, and sorbitol; starch from corn, wheat, rice, potato, or other plants; cellulose, such as methyl cellulose, hydroxypropylmethyl-cellulose, or sodium carboxymethylcellulose; gums, including arabic and tragacanth; and proteins, such as gelatin and collagen. If desired, disintegrating or solubilizing agents may be added, such as the cross-linked polyvinyl pyrrolidone, agar, and alginic acid or a salt thereof, such as sodium alginate.

Dragee cores may be used in conjunction with suitable coatings, such as concentrated sugar solutions, which may also contain gum arabic, talc, polyvinylpyrrolidone, carbopol gel, polyethylene glycol, and/or titanium dioxide, lacquer solutions, and suitable organic solvents or solvent mixtures. Dyestuffs or pigments may be added to the tablets or dragee coatings for product identification or to characterize the quantity of active compound, i.e., dosage.

Pharmaceutical preparations which can be used orally include push-fit capsules made of gelatin, as well as soft, sealed capsules made of gelatin and a coating, such as glycerol or sorbitol. Push-fit capsules can contain active ingredients mixed with fillers or binders, such as lactose or starches, lubricants, such as talc or magnesium stearate, and, optionally, stabilizers. In soft capsules, the active compounds may be dissolved or suspended in suitable liquids, such as fatty oils, liquid, or liquid polyethylene glycol with or without stabilizers.

Pharmaceutical formulations suitable for parenteral administration may be formulated in

aqueous solutions, preferably in physiologically compatible buffers such as Hanks' solution. Ringer's solution, or physiologically buffered saline. Aqueous injection suspensions may contain substances which increase the viscosity of the suspension, such as sodium carboxymethyl cellulose, sorbitol, or dextran. Additionally, suspensions of the active compounds may be prepared as appropriate oily injection suspensions. Suitable lipophilic solvents or vehicles include fatty oils, such as sesame oil, or synthetic fatty acid esters, such as ethyl oleate, triglycerides, or liposomes. Non-lipid polycationic amino polymers may also be used for delivery. Optionally, the suspension may also contain suitable stabilizers or agents to increase the solubility of the compounds and allow for the preparation of highly concentrated solutions.

For topical or nasal administration, penetrants appropriate to the particular barrier to be permeated are used in the formulation. Such penetrants are generally known in the art.

10

15

20

25

30

The pharmaceutical compositions of the present invention may be manufactured in a manner that is known in the art, e.g., by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping, or lyophilizing processes.

The pharmaceutical composition may be provided as a salt and can be formed with many acids, including but not limited to, hydrochloric, sulfuric, acetic, lactic, tartaric, malic, and succinic acids. Salts tend to be more soluble in aqueous or other protonic solvents than are the corresponding free base forms. In other cases, the preparation may be a lyophilized powder which may contain any or all of the following: I mM to 50 mM histidine, 0.1% to 2% sucrose, and 2% to 7% mannitol, at a pH range of 4.5 to 5.5, that is combined with buffer prior to use.

After pharmaceutical compositions have been prepared, they can be placed in an appropriate container and labeled for treatment of an indicated condition. For administration of NuABP, such labeling would include amount, frequency, and method of administration.

Pharmaceutical compositions suitable for use in the invention include compositions wherein the active ingredients are contained in an effective amount to achieve the intended purpose. The determination of an effective dose is well within the capability of those skilled in the art.

For any compound, the therapeutically effective dose can be estimated initially either in cell culture assays, e.g., of neoplastic cells, or in animal models such as mice, rats, rabbits, dogs, or pigs. An animal model may also be used to determine the appropriate concentration range and route of administration. Such information can then be used to determine useful doses and routes for administration in humans.

A therapeutically effective dose refers to that amount of active ingredient, for example NuABP or fragments thereof, antibodies of NuABP, and agonists, antagonists or inhibitors of NuABP, which ameliorates the symptoms or condition. Therapeutic efficacy and toxicity may be determined by standard pharmaceutical procedures in cell cultures or with experimental animals, such

as by calculating the ED_{30} (the dose therapeutically effective in 50% of the population) or LD_{30} (the dose lethal to 50% of the population) statistics. The dose ratio of toxic to therapeutic effects is the therapeutic index, which can be expressed as the LD_{30}/ED_{30} ratio. Pharmaceutical compositions which exhibit large therapeutic indices are preferred. The data obtained from cell culture assays and animal studies are used to formulate a range of dosage for human use. The dosage contained in such compositions is preferably within a range of circulating concentrations that includes the ED_{30} with little or no toxicity. The dosage varies within this range depending upon the dosage form employed, the sensitivity of the patient, and the route of administration.

The exact dosage will be determined by the practitioner, in light of factors related to the subject requiring treatment. Dosage and administration are adjusted to provide sufficient levels of the active moiety or to maintain the desired effect. Factors which may be taken into account include the severity of the disease state, the general health of the subject, the age, weight, and gender of the subject, time and frequency of administration, drug combination(s), reaction sensitivities, and response to therapy. Long-acting pharmaceutical compositions may be administered every 3 to 4 days, every week, or biweekly depending on the half-life and clearance rate of the particular formulation.

Normal dosage amounts may vary from about $0.1 \,\mu g$ to $100,000 \,\mu g$, up to a total dose of about 1 gram, depending upon the route of administration. Guidance as to particular dosages and methods of delivery is provided in the literature and generally available to practitioners in the art. Those skilled in the art will employ different formulations for nucleotides than for proteins or their inhibitors. Similarly, delivery of polynucleotides or polypeptides will be specific to particular cells, conditions, locations, etc.

DIAGNOSTICS

10

15

20

25

35

In another embodiment, antibodies which specifically bind NuABP may be used for the diagnosis of disorders characterized by expression of NuABP, or in assays to monitor patients being treated with NuABP or agonists, antagonists, or inhibitors of NuABP. Antibodies useful for diagnostic purposes may be prepared in the same manner as described above for therapeutics. Diagnostic assays for NuABP include methods which utilize the antibody and a label to detect NuABP in human body fluids or in extracts of cells or tissues. The antibodies may be used with or without modification, and may be labeled by covalent or non-covalent attachment of a reporter molecule. A wide variety of reporter molecules, several of which are described above, are known in the art and may be used.

A variety of protocols for measuring NuABP, including ELISAs, RIAs, and FACS, are known in the art and provide a basis for diagnosing altered or abnormal levels of NuABP expression.

Normal or standard values for NuABP expression are established by combining body fluids or cell

extracts taken from normal mammalian subjects, for example, human subjects, with antibody to NuABP under conditions suitable for complex formation. The amount of standard complex formation may be quantitated by various methods, such as photometric means. Quantities of NuABP expressed in subject, control, and disease samples from biopsied tissues are compared with the standard values. Deviation between standard and subject values establishes the parameters for diagnosing disease.

In another embodiment of the invention, the polynucleotides encoding NuABP may be used for diagnostic purposes. The polynucleotides which may be used include oligonucleotide sequences, complementary RNA and DNA molecules, and PNAs. The polynucleotides may be used to detect and quantify gene expression in biopsied tissues in which expression of NuABP may be correlated with disease. The diagnostic assay may be used to determine absence, presence, and excess expression of NuABP, and to monitor regulation of NuABP levels during therapeutic intervention.

10

20

25

30

35

In one aspect, hybridization with PCR probes which are capable of detecting polynucleotide sequences, including genomic sequences, encoding NuABP or closely related molecules may be used to identify nucleic acid sequences which encode NuABP. The specificity of the probe, whether it is made from a highly specific region, e.g., the 5' regulatory region, or from a less specific region, e.g., a conserved motif, and the stringency of the hybridization or amplification will determine whether the probe identifies only naturally occurring sequences encoding NuABP, allelic variants, or related sequences.

Probes may also be used for the detection of related sequences, and may have at least 50% sequence identity to any of the NuABP encoding sequences. The hybridization probes of the subject invention may be DNA or RNA and may be derived from the sequence of SEQ ID NO:56-110 or from genomic sequences including promoters, enhancers, and introns of the NuABP gene.

Means for producing specific hybridization probes for DNAs encoding NuABP include the cloning of polynucleotide sequences encoding NuABP or NuABP derivatives into vectors for the production of mRNA probes. Such vectors are known in the art, are commercially available, and may be used to synthesize RNA probes in vitro by means of the addition of the appropriate RNA polymerases and the appropriate labeled nucleotides. Hybridization probes may be labeled by a variety of reporter groups, for example, by radionuclides such as ³²P or ³⁵S, or by enzymatic labels, such as alkaline phosphatase coupled to the probe via avidin/biotin coupling systems, and the like.

Polynucleotide sequences encoding NuABP may be used for the diagnosis of disorders associated with expression of NuABP. Examples of such disorders include, but are not limited to, a reproductive disorder such as disorders of prolactin production, infertility, including tubal disease, ovulatory defects, and endometriosis, disruptions of the estrous cycle, disruptions of the menstrual cycle, polycystic ovary syndrome, ovarian hyperstimulation syndrome, endometrial and ovarian tumors, uterine fibroids, autoimmune disorders, ectopic pregnancies, and teratogenesis; cancer of the

PCT/US00/02237 WO 00/44900

breast, fibrocystic breast disease, and galactorrhea; disruptions of spermatogenesis, abnormal sperm physiology, cancer of the testis, cancer of the prostate, benign prostatic hyperplasia, prostatitis, Peyronie's disease, impotence, carcinoma of the male breast, and gynecomastia: an immune disorder such as inflammation, actinic keratosis, acquired immunodeficiency syndrome (AIDS), Addison's disease, adult respiratory distress syndrome, allergies, ankylosing spondylitis, amyloidosis, anemia, arteriosclerosis, asthma, atherosclerosis, autoimmune hemolytic anemia, autoimmune thyroiditis, bronchitis, bursitis, cholecystitis, cirrhosis, contact dermatitis, Crohn's disease, atopic dermatitis, dermatomyositis, diabetes mellitus, emphysema, erythroblastosis fetalis, erythema nodosum, atrophic gastritis, glomerulonephritis, Goodpasture's syndrome, gout, Graves' disease, Hashimoto's thyroiditis, paroxysmal nocturnal hemoglobinuria, hepatitis, hypereosinophilia, irritable bowel syndrome, episodic lymphopenia with lymphocytotoxins, mixed connective tissue disease (MCTD), multiple sclerosis, myasthenia gravis, myocardial or pericardial inflammation, myelofibrosis, osteoarthritis, osteoporosis, pancreatitis, polycythemia vera, polymyositis, psoriasis, Reiter's syndrome, rheumatoid arthritis, scleroderma, Sjögren's syndrome, systemic anaphylaxis, systemic lupus erythematosus, systemic sclerosis, primary thrombocythemia, thrombocytopenic purpura, ulcerative colitis, uveitis, Werner syndrome, complications of cancer, hemodialysis, and extracorporeal circulation, trauma, and hematopoietic cancer including lymphoma, leukemia. and myeloma; a neurological disorder such as epilepsy, ischemic cerebrovascular disease, stroke. cerebral neoplasms, Alzheimer's disease, Pick's disease, Huntington's disease, dementia, Parkinson's disease and other extrapyramidal disorders, amyotrophic lateral sclerosis and other motor neuron disorders, progressive neural muscular atrophy, retinitis pigmentosa, hereditary ataxias, multiple sclerosis and other demyelinating diseases, bacterial and viral meningitis, brain abscess, subdural empyema, epidural abscess, suppurative intracranial thrombophlebitis, myelitis and radiculitis, viral central nervous system disease, prion diseases including kuru, Creutzfeldt-Jakob disease, and Gerstmann-Straussler-Scheinker syndrome, fatal familial insomnia, nutritional and metabolic diseases of the 25 nervous system, neurofibromatosis, tuberous sclerosis, cerebelloretinal hemangioblastomatosis, encephalotrigeminal syndrome, mental retardation and other developmental disorders of the central nervous system, cerebral palsy, neuroskeletal disorders, autonomic nervous system disorders, cranial nerve disorders, spinal cord diseases, muscular dystrophy and other neuromuscular disorders, peripheral nervous system disorders, dermatomyositis and polymyositis, inherited, metabolic, 30 endocrine, and toxic myopathies, myasthenia gravis, periodic paralysis, mental disorders including mood, anxiety, and schizophrenic disorders, akathesia, amnesia, catatonia, diabetic neuropathy, tardive dyskinesia, dystonias, paranoid psychoses, postherpetic neuralgia, and Tourette's disorder; and a cell proliferative disorder such as actinic keratosis, arteriosclerosis, atherosclerosis, bursitis, cirrhosis, hepatitis, mixed connective tissue disease (MCTD), myelofibrosis, paroxysmal nocturnal 35

10

20

hemoglobinuria, polycythemia vera, psoriasis, primary thrombocythemia, and cancers including adenocarcinoma, leukemia, lymphoma, melanoma, myeloma, sarcoma, teratocarcinoma, and, in particular, cancers of the adrenal gland, bladder, bone, bone marrow, brain, breast, cervix, gall bladder, ganglia, gastrointestinal tract, heart, kidney, liver, lung, muscle, ovary, pancreas, parathyroid, 5 penis, prostate, salivary glands, skin, spleen, testis, thymus, thyroid, and uterus. The polynucleotide sequences encoding NuABP may be used in Southern or northern analysis, dot blot, or other membrane-based technologies; in PCR technologies; in dipstick, pin, and multiformat ELISA-like assays; and in microarrays utilizing fluids or tissues from patients to detect altered NuABP expression. Such qualitative or quantitative methods are well known in the art.

In a particular aspect, the nucleotide sequences encoding NuABP may be useful in assays that detect the presence of associated disorders, particularly those mentioned above. The nucleotide sequences encoding NuABP may be labeled by standard methods and added to a fluid or tissue sample from a patient under conditions suitable for the formation of hybridization complexes. After a suitable incubation period, the sample is washed and the signal is quantified and compared with a 15 standard value. If the amount of signal in the patient sample is significantly altered in comparison to a control sample then the presence of altered levels of nucleotide sequences encoding NuABP in the sample indicates the presence of the associated disorder. Such assays may also be used to evaluate the efficacy of a particular therapeutic treatment regimen in animal studies, in clinical trials, or to monitor the treatment of an individual patient.

10

20

In order to provide a basis for the diagnosis of a disorder associated with expression of NuABP, a normal or standard profile for expression is established. This may be accomplished by combining body fluids or cell extracts taken from normal subjects, either animal or human, with a sequence, or a fragment thereof, encoding NuABP, under conditions suitable for hybridization or amplification. Standard hybridization may be quantified by comparing the values obtained from normal subjects with values from an experiment in which a known amount of a substantially purified polynucleotide is used. Standard values obtained in this manner may be compared with values obtained from samples from patients who are symptomatic for a disorder. Deviation from standard values is used to establish the presence of a disorder.

Once the presence of a disorder is established and a treatment protocol is initiated, hybridization assays may be repeated on a regular basis to determine if the level of expression in the patient begins to approximate that which is observed in the normal subject. The results obtained from successive assays may be used to show the efficacy of treatment over a period ranging from several days to months.

With respect to cancer, the presence of an abnormal amount of transcript (either under- or overexpressed) in biopsied tissue from an individual may indicate a predisposition for the

was proper from the state of the state of

development of the disease, or may provide a means for detecting the disease prior to the appearance of actual clinical symptoms. A more definitive diagnosis of this type may allow health professionals to employ preventative measures or aggressive treatment earlier thereby preventing the development or further progression of the cancer.

5

10

15

20

25

30

Additional diagnostic uses for oligonucleotides designed from the sequences encoding NuABP may involve the use of PCR. These oligomers may be chemically synthesized, generated enzymatically, or produced in vitro. Oligomers will preferably contain a fragment of a polynucleotide encoding NuABP, or a fragment of a polynucleotide complementary to the polynucleotide encoding NuABP, and will be employed under optimized conditions for identification of a specific gene or condition. Oligomers may also be employed under less stringent conditions for detection or quantification of closely related DNA or RNA sequences.

Methods which may also be used to quantify the expression of NuABP include radiolabeling or biotinylating nucleotides, coamplification of a control nucleic acid, and interpolating results from standard curves. (See, e.g., Melby, P.C. et al. (1993) J. Immunol. Methods 159:235-244; Duplaa, C. et al. (1993) Anal. Biochem. 212:229-236.) The speed of quantitation of multiple samples may be accelerated by running the assay in a high-throughput format where the oligomer of interest is presented in various dilutions and a spectrophotometric or colorimetric response gives rapid quantitation.

In further embodiments, oligonucleotides or longer fragments derived from any of the polynucleotide sequences described herein may be used as targets in a microarray. The microarray can be used to monitor the expression level of large numbers of genes simultaneously and to identify genetic variants, mutations, and polymorphisms. This information may be used to determine gene function, to understand the genetic basis of a disorder, to diagnose a disorder, and to develop and monitor the activities of therapeutic agents.

Microarrays may be prepared, used, and analyzed using methods known in the art. (See, e.g., Brennan, T.M. et al. (1995) U.S. Patent No. 5,474,796; Schena, M. et al. (1996) Proc. Natl. Acad. Sci. USA 93:10614-10619; Baldeschweiler et al. (1995) PCT application WO95/251116; Shalon, D. et al. (1995) PCT application WO95/35505; Heller, R.A. et al. (1997) Proc. Natl. Acad. Sci. USA 94:2150-2155; and Heller, M.J. et al. (1997) U.S. Patent No. 5,605,662.)

In another embodiment of the invention, nucleic acid sequences encoding NuABP may be used to generate hybridization probes useful in mapping the naturally occurring genomic sequence. The sequences may be mapped to a particular chromosome, to a specific region of a chromosome, or to artificial chromosome constructions, e.g., human artificial chromosomes (HACs), yeast artificial chromosomes (YACs), bacterial artificial chromosomes (BACs), bacterial P1 constructions, or single chromosome cDNA libraries. (See, e.g., Harrington, J.J. et al. (1997) Nat. Genet. 15:345-355; Price.

C.M. (1993) Blood Rev. 7:127-134; and Trask, B.J. (1991) Trends Genet. 7:149-154.)

10

20

25

35

Fluorescent in situ hybridization (FISH) may be correlated with other physical chromosome mapping techniques and genetic map data. (See. e.g., Heinz-Ulrich, et al. (1995) in Meyers, supra, pp. 965-968.) Examples of genetic map data can be found in various scientific journals or at the Online Mendelian Inheritance in Man (OMIM) World Wide Web site. Correlation between the location of the gene encoding NuABP on a physical chromosomal map and a specific disorder, or a predisposition to a specific disorder, may help define the region of DNA associated with that disorder. The nucleotide sequences of the invention may be used to detect differences in gene sequences among normal, carrier, and affected individuals.

In situ hybridization of chromosomal preparations and physical mapping techniques, such as linkage analysis using established chromosomal markers, may be used for extending genetic maps. Often the placement of a gene on the chromosome of another mammalian species, such as mouse, may reveal associated markers even if the number or arm of a particular human chromosome is not known. New sequences can be assigned to chromosomal arms by physical mapping. This provides valuable information to investigators searching for disease genes using positional cloning or other gene discovery techniques. Once the disease or syndrome has been crudely localized by genetic linkage to a particular genomic region, e.g., ataxia-telangiectasia to 11q22-23, any sequences mapping to that area may represent associated or regulatory genes for further investigation. (See, e.g., Gatti, R.A. et al. (1988) Nature 336:577-580.) The nucleotide sequence of the subject invention may also be used to detect differences in the chromosomal location due to translocation, inversion, etc., among normal, earrier, or affected individuals.

In another embodiment of the invention, NuABP, its catalytic or immunogenic fragments, or oligopeptides thereof can be used for screening libraries of compounds in any of a variety of drug screening techniques. The fragment employed in such screening may be free in solution, affixed to a solid support, borne on a cell surface, or located intracellularly. The formation of binding complexes between NuABP and the agent being tested may be measured.

Another technique for drug screening provides for high throughput screening of compounds having suitable binding affinity to the protein of interest. (See, e.g., Geysen, et al. (1984) PCT application WO84/03564.) In this method, large numbers of different small test compounds are synthesized on a solid substrate. The test compounds are reacted with NuABP, or fragments thereof, and washed. Bound NuABP is then detected by methods well known in the art. Purified NuABP can also be coated directly onto plates for use in the aforementioned drug screening techniques.

Alternatively, non-neutralizing antibodies can be used to capture the peptide and immobilize it on a solid support.

In another embodiment, one may use competitive drug screening assays in which neutralizing

antibodies capable of binding NuABP specifically compete with a test compound for binding NuABP. In this manner, antibodies can be used to detect the presence of any peptide which shares one or more antigenic determinants with NuABP.

In additional embodiments, the nucleotide sequences which encode NuABP may be used in any molecular biology techniques that have yet to be developed, provided the new techniques rely on properties of nucleotide sequences that are currently known, including, but not limited to, such properties as the triplet genetic code and specific base pair interactions.

Without further elaboration, it is believed that one skilled in the art can, using the preceding description, utilize the present invention to its fullest extent. The following embodiments are, therefore, to be construed as merely illustrative, and not limitative of the remainder of the disclosure in any way whatsoever

Without further elaboration, it is believed that one skilled in the art can, using the preceding description, utilize the present invention to its fullest extent. The following preferred specific embodiments are, therefore, to be construed as merely illustrative, and not limitative of the remainder of the disclosure in any way whatsoever.

The disclosures of all patents, applications, and publications mentioned above and below, in particular U.S. Ser. No. 60/117,905 and U.S. Ser. No. 60/117,904, are hereby expressly incorporated by reference.

20 EXAMPLES

I. Construction of cDNA Libraries

15

25

3.5

RNA was purchased from Clontech or isolated from tissues described in Table 4. Some tissues were homogenized and lysed in guanidinium isothiocyanate, while others were homogenized and lysed in phenol or in a suitable mixture of denaturants, such as TRIZOL (Life Technologies), a monophasic solution of phenol and guanidine isothiocyanate. The resulting lysates were centrifuged over CsCl cushions or extracted with chloroform. RNA was precipitated from the lysates with either isopropanol or sodium acetate and ethanol, or by other routine methods.

Phenol extraction and precipitation of RNA were repeated as necessary to increase RNA purity. In some cases, RNA was treated with DNase. For most libraries, poly(A+) RNA was isolated using oligo d(T)-coupled paramagnetic particles (Promega), OLIGOTEX latex particles (QIAGEN, Chatsworth CA), or an OLIGOTEX mRNA purification kit (QIAGEN). Alternatively, RNA was isolated directly from tissue lysates using other RNA isolation kits, e.g., the POLY(A)PURE mRNA purification kit (Ambion, Austin TX).

In some cases, Stratagene was provided with RNA and constructed the corresponding cDNA libraries. Otherwise, cDNA was synthesized and cDNA libraries were constructed with the UNIZAP

vector system (Stratagene) or SUPERSCRIPT plasmid system (Life Technologies), using the recommended procedures or similar methods known in the art. (See, e.g., Ausubel, 1997, supra, units 5.1-6.6.) Reverse transcription was initiated using oligo d(T) or random primers. Synthetic oligonucleotide adapters were ligated to double stranded cDNA, and the cDNA was digested with the appropriate restriction enzyme or enzymes. For most libraries, the cDNA was size-selected (300-1000 bp) using SEPHACRYL \$1000, SEPHAROSE CL2B, or SEPHAROSE CL4B column chromatography (Amersham Pharmacia Biotech) or preparative agarose gel electrophoresis. cDNAs were ligated into compatible restriction enzyme sites of the polylinker of a suitable plasmid, e.g., PBLUESCRIPT plasmid (Stratagene), PSPORT1 plasmid (Life Technologies), or pINCY (Incyte Pharmaceuticals, Palo Alto CA). Recombinant plasmids were transformed into competent E. coli cells including XL1-Blue, XL1-BlueMRF, or SOLR from Stratagene or DH5 α , DH10B, or ElectroMAX DH10B from Life Technologies.

II. Isolation of cDNA Clones

15

20

25

35

Plasmids were recovered from host cells by in vivo excision using the UNIZAP vector system (Stratagene) or by cell lysis. Plasmids were purified using at least one of the following: a Magic or WIZARD Minipreps DNA purification system (Promega); an AGTC Miniprep purification kit (Edge Biosystems, Gaithersburg MD); and QIAWELL 8 Plasmid, QIAWELL 8 Plasmid purification systems or the R.E.A.L. PREP 96 plasmid purification kit from QIAGEN. Following precipitation, plasmids were resuspended in 0.1 ml of distilled water and stored, with or without Ivophilization at 4°C.

Alternatively, plasmid DNA was amplified from host cell lysates using direct link PCR in a high-throughput format (Rao, V.B. (1994) Anal. Biochem. 216:1-14). Host cell lysis and thermal cycling steps were carried out in a single reaction mixture. Samples were processed and stored in 384-well plates, and the concentration of amplified plasmid DNA was quantified fluorometrically using PICOGREEN dye (Molecular Probes, Eugene OR) and a FLUOROSKAN II fluorescence scanner (Labsystems Oy, Helsinki, Finland).

III. Sequencing and Analysis

cDNA sequencing reactions were processed using standard methods or high-throughput instrumentation such as the ABI CATALYST 800 (Perkin-Elmer) thermal cycler or the PTC-200 thermal cycler (MJ Research) in conjunction with the HYDRA microdispenser (Robbins Scientific) or the MICROLAB 2200 (Hamilton) liquid transfer system. cDNA sequencing reactions were prepared using reagents provided by Amersham Pharmacia Biotech or supplied in ABI sequencing kits such as the ABI PRISM BIGDYE Terminator cycle sequencing ready reaction kit (Perkin-Elmer). Electrophoretic separation of cDNA sequencing reactions and detection of labeled polynucleotides were carried out using the MEGABACE 1000 DNA sequencing system (Molecular Dynamics); the

ABI PRISM 373 or 377 sequencing system (Perkin-Elmer) in conjunction with standard ABI protocols and base calling software: or other sequence analysis systems known in the art. Reading frames within the cDNA sequences were identified using standard methods (reviewed in Ausubel, 1997, https://gwpra.unit.77). Some of the cDNA sequences were selected for extension using the techniques disclosed in Example V.

The polynucleotide sequences derived from cDNA sequencing were assembled and analyzed using a combination of software programs which utilize algorithms well known to those skilled in the art. Table 5 summarizes the tools, programs, and algorithms used and provides applicable descriptions, references, and threshold parameters. The first column of Table 5 shows the tools, programs, and algorithms used, the second column provides brief descriptions thereof, the third column presents appropriate references, all of which are incorporated by reference herein in their entirety, and the fourth column presents, where applicable, the scores, probability values, and other parameters used to evaluate the strength of a match between two sequences (the higher the score, the greater the homology between two sequences). Sequences were analyzed using MACDNASIS PRO software (Hitachi Software Engineering, South San Francisco CA) and LASERGENE software (DNASTAR). Polynucleotide and polypeptide sequence alignments were generated using the default parameters specified by the clustal algorithm as incorporated into the MEGALIGN multisequence alignment program (DNASTAR), which also calculates the percent identity between aligned sequences.

10

15

20

25

30

The polynucleotide sequences were validated by removing vector, linker, and polyA sequences and by masking ambiguous bases, using algorithms and programs based on BLAST, dynamic programing, and dinucleotide nearest neighbor analysis. The sequences were then queried against a selection of public databases such as the GenBank primate, rodent, mammalian, vertebrate, and eukaryote databases, and BLOCKS, PRINTS, DOMO, PRODOM, and PFAM to acquire annotation using programs based on BLAST, FASTA, and BLIMPS. The sequences were assembled into full length polynucleotide sequences using programs based on Phred, Phrap, and Consed, and were screened for open reading frames using programs based on GeneMark, BLAST, and FASTA. The full length polynucleotide sequences were translated to derive the corresponding full length amino acid sequences, and these full length sequences were subsequently analyzed by querying against databases such as the GenBank databases (described above), SwissProt, BLOCKS, PRINTS, DOMO, PRODOM, Prosite, and Hidden Markov Model (HMM)-based protein family databases such as PFAM. HMM is a probabilistic approach which analyzes consensus primary structures of gene families. (See, e.g., Eddy, S.R. (1996) Curr. Opin. Struct. Biol. 6:361-365.)

The programs described above for the assembly and analysis of full length polynucleotide and
35 amino acid sequences were also used to identify polynucleotide sequence fragments from SEQ ID

NO:56-110. Fragments from about 20 to about 4000 nucleotides which are useful in hybridization and amplification technologies were described in The Invention section above.

IV. Northern Analysis

20

30

Northern analysis is a laboratory technique used to detect the presence of a transcript of a gene and involves the hybridization of a labeled nucleotide sequence to a membrane on which RNAs from a particular cell type or tissue have been bound. (See, e.g., Sambrook, <u>supra</u>, ch. 7; Ausubel, 1995, supra, ch. 4 and 16.)

Analogous computer techniques applying BLAST were used to search for identical or related molecules in nucleotide databases such as GenBank or LIFESEQ (Incyte Pharmaceuticals). This analysis is much faster than multiple membrane-based hybridizations. In addition, the sensitivity of the computer search can be modified to determine whether any particular match is categorized as exact or similar. The basis of the search is the product score, which is defined as:

% sequence identity x % maximum BLAST score

100

15 The product score takes into account both the degree of similarity between two sequences and the length of the sequence match. For example, with a product score of 40, the match will be exact within a 1% to 2% error, and, with a product score of 70, the match will be exact. Similar molecules are usually identified by selecting those which show product scores between 15 and 40, although lower scores may identify related molecules.

The results of northern analyses are reported as a percentage distribution of libraries in which the transcript encoding NuABP occurred. Analysis involved the categorization of cDNA libraries by organ/tissue and disease. The organ/tissue categories included cardiovascular, dermatologic, developmental, endocrine, gastrointestinal, hematopoietic/immune, musculoskeletal, nervous, reproductive, and urologic. The disease/condition categories included cancer, inflammation, trauma, cell proliferation, neurological, and pooled. For each category, the number of libraries expressing the sequence of interest was counted and divided by the total number of libraries across all categories. Percentage values of tissue-specific and disease- or condition-specific expression are reported in Table 3.

V. Extension of NuABP Encoding Polynucleotides

The full length nucleic acid sequences of SEQ ID NO:56-110 were produced by extension of an appropriate fragment of the full length molecule using oligonucleotide primers designed from this fragment. One primer was synthesized to initiate 5' extension of the known fragment, and the other primer, to initiate 3' extension of the known fragment. The initial primers were designed using OLIGO 4.06 software (National Biosciences), or another appropriate program, to be about 22 to 30 nucleotides in length, to have a GC content of about 50% or more, and to anneal to the target

sequence at temperatures of about 68°C to about 72°C. Any stretch of nucleotides which would result in hairpin structures and primer-primer dimerizations was avoided.

Selected human cDNA libraries were used to extend the sequence. If more than one extension was necessary or desired, additional or nested sets of primers were designed.

5

15

35

High fidelity amplification was obtained by PCR using methods well known in the art. PCR was performed in 96-well plates using the PTC-200 thermal cycler (MJ Research, Inc.). The reaction mix contained DNA template, 200 nmol of each primer, reaction buffer containing Mg^{2-*}. (NH₄)₂SO₄, and β-mercaptoethanol, Taq DNA polymerase (Amersham Pharmacia Biotech), ELONGASE enzyme (Life Technologies), and Pfu DNA polymerase (Stratagene), with the following parameters for primer pair PCI A and PCI B: Step 1: 94°C, 3 min; Step 2: 94°C, 15 sec; Step 3: 60°C, 1 min: Step 4: 68°C, 2 min; Step 5: Steps 2, 3, and 4 repeated 20 times; Step 6: 68°C, 5 min; Step 1: 94°C, 3 min; Step 2: 94°C, 15 sec; Step 3: 57°C, 1 min; Step 4: 68°C, 2 min; Step 5: Steps 2, 3, and 4 repeated 20 times; Step 6: 68°C, 5 min; Step 7: storage at 4°C.

The concentration of DNA in each well was determined by dispensing 100 µl PICOGREEN quantitation reagent (0.25% (v/v) PICOGREEN; Molecular Probes, Eugene OR) dissolved in 1X TE and 0.5 µl of undiluted PCR product into each well of an opaque fluorimeter plate (Corning Costar, Acton MA), allowing the DNA to bind to the reagent. The plate was scanned in a Fluoroskan II (Labsystems Oy, Helsinki, Finland) to measure the fluorescence of the sample and to quantify the concentration of DNA. A 5 µl to 10 µl aliquot of the reaction mixture was analyzed by electrophoresis on a 1 % agarose mini-gel to determine which reactions were successful in extending the sequence.

The extended nucleotides were desalted and concentrated, transferred to 384-well plates, digested with CviJI cholera virus endonuclease (Molecular Biology Research, Madison WI), and sonicated or sheared prior to religation into pUC 18 vector (Amersham Pharmacia Biotech). For shotgun sequencing, the digested nucleotides were separated on low concentration (0.6 to 0.8%) agarose gels, fragments were excised, and agar digested with Agar ACE (Promega). Extended clones were religated using T4 ligase (New England Biolabs, Beverly MA) into pUC 18 vector (Amersham Pharmacia Biotech), treated with Pfu DNA polymerase (Stratagene) to fill-in restriction site overhangs, and transfected into competent <u>E. coli</u> cells. Transformed cells were selected on antibiotic-containing media, individual colonies were picked and cultured overnight at 37°C in 384-well plates in LB/2x carb liquid media.

The cells were lysed, and DNA was amplified by PCR using Taq DNA polymerase (Amersham Pharmacia Biotech) and Pfu DNA polymerase (Stratagene) with the following parameters: Step 1: 94°C, 3 min; Step 2: 94°C, 15 sec; Step 3: 60°C, 1 min; Step 4: 72°C, 2 min;

Step 5: steps 2, 3, and 4 repeated 29 times; Step 6: 72°C, 5 min; Step 7: storage at 4°C. DNA was quantified by PICOGREEN reagent (Molecular Probes) as described above. Samples with low DNA recoveries were reamplified using the same conditions as described above. Samples were diluted with 20% dimethysulfoxide (1:2, v/v), and sequenced using DYENAMIC energy transfer sequencing primers and the DYENAMIC DIRECT kit (Amersham Pharmacia Biotech) or the ABI PRISM BIGDYE Terminator cycle sequencing ready reaction kit (Perkin-Elmer).

In like manner, the nucleotide sequences of SEQ ID NO:56-110 are used to obtain 5' regulatory sequences using the procedure above, oligonucleotides designed for such extension, and an appropriate genomic library.

10 VI. Labeling and Use of Individual Hybridization Probes

Hybridization probes derived from SEQ ID NO:56-110 are employed to screen cDNAs, genomic DNAs, or mRNAs. Although the labeling of oligonucleotides, consisting of about 20 base pairs, is specifically described, essentially the same procedure is used with larger nucleotide fragments. Oligonucleotides are designed using state-of-the-art software such as OLIGO 4.06 software (National Biosciences) and labeled by combining 50 pmol of each oligomer, 250 μ Ci of [γ - 12 P] adenosine triphosphate (Amersham Pharmacia Biotech), and T4 polynucleotide kinase (DuPont NEN, Boston MA). The labeled oligonucleotides are substantially purified using a SEPHADEX G-25 superfine size exclusion dextran bead column (Amersham Pharmacia Biotech). An aliquot containing 10° counts per minute of the labeled probe is used in a typical membrane-based hybridization analysis of human genomic DNA digested with one of the following endonucleases: Ase I. Bgl II. Eco RI, PSt I, Xba I, or Pvu II (DuPont NEN).

The DNA from each digest is fractionated on a 0.7% agarose gel and transferred to nylon membranes (Nytran Plus, Schleicher & Schuell, Durham NH). Hybridization is carried out for 16 hours at 40 °C. To remove nonspecific signals, blots are sequentially washed at room temperature under conditions of up to, for example, 0.1 x saline sodium citrate and 0.5% sodium dodecyl sulfate. Hybridization patterns are visualized using autoradiography or an alternative imaging means and compared.

VII. Microarrays

25

30

35

A chemical coupling procedure and an ink jet device can be used to synthesize array elements on the surface of a substrate. (See, e.g., Baldeschweiler, supra. An array analogous to a dot or slot blot may also be used to arrange and link elements to the surface of a substrate using thermal, UV, chemical, or mechanical bonding procedures. A typical array may be produced by hand or using available methods and machines and contain any appropriate number of elements. After hybridization, nonhybridized probes are removed and a scanner used to determine the levels and patterns of fluorescence. The degree of complementarity and the relative abundance of each probe

on a course building from paragraph of the same of the course of the same of the course of the cours

which hybridizes to an element on the microarray may be assessed through analysis of the scanned images.

Full-length cDNAs, Expressed Sequence Tags (ESTs). or fragments thereof may comprise the elements of the microarray. Fragments suitable for hybridization can be selected using software well known in the art such as LASERGENE software (DNASTAR). Full-length cDNAs, ESTs, or fragments thereof corresponding to one of the nucleotide sequences of the present invention, or selected at random from a cDNA library relevant to the present invention, are arranged on an appropriate substrate, e.g., a glass slide. The cDNA is fixed to the slide using, e.g., UV cross-linking followed by thermal and chemical treatments and subsequent drying. (See, e.g., Schena, M. et al. (1995) Science 270:467-470; Shalon, D. et al. (1996) Genome Res. 6:639-645.) Fluorescent probes are prepared and used for hybridization to the elements on the substrate. The substrate is analyzed by procedures described above.

VIII. Complementary Polynucleotides

Sequences complementary to the NuABP-encoding sequences, or any parts thereof, are used to detect, decrease, or inhibit expression of naturally occurring NuABP. Although use of oligonucleotides comprising from about 15 to 30 base pairs is described, essentially the same procedure is used with smaller or with larger sequence fragments. Appropriate oligonucleotides are designed using OLIGO 4.06 software (National Biosciences) and the coding sequence of NuABP. To inhibit transcription, a complementary oligonucleotide is designed from the most unique 5' sequence and used to prevent promoter binding to the coding sequence. To inhibit translation, a complementary oligonucleotide is designed to prevent ribosomal binding to the NuABP-encoding transcript.

IX. Expression of NuABP

N. V. Stade Communication of April 1984 agreement at a

20

Expression and purification of NuABP is achieved using bacterial or virus-based expression systems. For expression of NuABP in bacteria, cDNA is subcloned into an appropriate vector containing an antibiotic resistance gene and an inducible promoter that directs high levels of cDNA transcription. Examples of such promoters include, but are not limited to, the *trp-lac (tac)* hybrid promoter and the T5 or T7 bacteriophage promoter in conjunction with the *lac* operator regulatory element. Recombinant vectors are transformed into suitable bacterial hosts, e.g., BL21(DE3). Antibiotic resistant bacteria express NuABP upon induction with isopropyl beta-D-thiogalactopyranoside (IPTG). Expression of NuABP in eukaryotic cells is achieved by infecting insect or mammalian cell lines with recombinant <u>Autographica californica</u> nuclear polyhedrosis virus (AcMNPV), commonly known as baculovirus. The nonessential polyhedrin gene of baculovirus is replaced with cDNA encoding NuABP by either homologous recombination or bacterial-mediated transposition involving transfer plasmid intermediates. Viral infectivity is maintained and the strong polyhedrin promoter drives high levels of cDNA transcription. Recombinant baculovirus is used to

infect Spodoptera frugiperda (Sf9) insect cells in most cases, or human hepatocytes, in some cases. Infection of the latter requires additional genetic modifications to baculovirus. (See Engelhard, E.K. et al. (1994) Proc. Natl. Acad. Sci. USA 91:3224-3227; Sandig, V. et al. (1996) Hum. Gene Ther. 7:1937-1945.)

In most expression systems, NuABP is synthesized as a fusion protein with, e.g., glutathione S-transferase (GST) or a peptide epitope tag, such as FLAG or 6-His, permitting rapid, single-step, affinity-based purification of recombinant fusion protein from crude cell lysates. GST, a 26-kilodalton enzyme from Schistosoma japonicum, enables the purification of fusion proteins on immobilized glutathione under conditions that maintain protein activity and antigenicity (Amersham Pharmacia Biotech). Following purification, the GST moiety can be proteolytically cleaved from NuABP at specifically engineered sites. FLAG, an 8-amino acid peptide, enables immunoaffinity purification using commercially available monoclonal and polyclonal anti-FLAG antibodies (Eastman Kodak). 6-His, a stretch of six consecutive histidine residues, enables purification on metal-chelate resins (QIAGEN). Methods for protein expression and purification are discussed in Ausubel (1995, supra, ch. 10 and 16). Purified NuABP obtained by these methods can be used directly in the following activity assay.

X. Demonstration of NuABP Activity

10

20

25

30

35

NuABP activity is measured by its ability to stimulate transcription of a reporter gene (Liu, H.Y. et al. (1997) EMBO J. 16(17):5289-5298.) The assay entails the use of a well characterized reporter gene construct, LexA_{op}-LacZ, that consists of LexA DNA transcriptional control elements (LexA_{op}) fused to sequences encoding the <u>E. coli</u> LacZ enzyme. The methods for constructing and expressing fusions genes, introducing them into cells, and measuring LacZ enzyme activity, are well known to those skilled in the art. Sequences encoding NuABP are cloned into a plasmid that directs the synthesis of a fusion protein, LexA-NuABP, consisting of NuABP and a DNA binding domain derived from the LexA transcription factor. The resulting plasmid, encoding a LexA-NuABP fusion protein, is introduced into yeast cells along with a plasmid containing the LexA_{op}-LacZ reporter gene. The amount of LacZ enzyme activity associated with LexA-NuABP transfected cells, relative to control cells, is proportional to the amount of transcription stimulated by the NuABP.

XI. Functional Assays

NuABP function is assessed by expressing the sequences encoding NuABP at physiologically elevated levels in mammalian cell culture systems. cDNA is subcloned into a mammalian expression vector containing a strong promoter that drives high levels of cDNA expression. Vectors of choice include pCMV SPORT (Life Technologies) and pCR3.1 (Invitrogen, Carlsbad CA), both of which contain the cytomegalovirus promoter. 5-10 μ g of recombinant vector are transiently transfected into a human cell line, for example, an endothelial or hematopoietic cell

line, using either liposome formulations or electroporation. 1-2 µg of an additional plasmid containing sequences encoding a marker protein are co-transfected. Expression of a marker protein provides a means to distinguish transfected cells from nontransfected cells and is a reliable predictor of cDNA expression from the recombinant vector. Marker proteins of choice include, e.g., Green 5 Fluorescent Protein (GFP; Clontech), CD64, or a CD64-GFP fusion protein. Flow cytometry (FCM), an automated, laser optics-based technique, is used to identify transfected cells expressing GFP or CD64-GFP and to evaluate the apoptotic state of the cells and other cellular properties. FCM detects and quantifies the uptake of fluorescent molecules that diagnose events preceding or coincident with cell death. These events include changes in nuclear DNA content as measured by staining of DNA 10 with propidium iodide; changes in cell size and granularity as measured by forward light scatter and 90 degree side light scatter; down-regulation of DNA synthesis as measured by decrease in bromodeoxyuridine uptake: alterations in expression of cell surface and intracellular proteins as measured by reactivity with specific antibodies; and alterations in plasma membrane composition as measured by the binding of fluorescein-conjugated Annexin V protein to the cell surface. Methods in 15 flow cytometry are discussed in Ormerod, M.G. (1994) Flow Cytometry, Oxford, New York NY.

The influence of NuABP on gene expression can be assessed using highly purified populations of cells transfected with sequences encoding NuABP and either CD64 or CD64-GFP. CD64 and CD64-GFP are expressed on the surface of transfected cells and bind to conserved regions of human immunoglobulin G (IgG). Transfected cells are efficiently separated from nontransfected cells using magnetic beads coated with either human IgG or antibody against CD64 (DYNAL, Lake Success NY). mRNA can be purified from the cells using methods well known by those of skill in the art. Expression of mRNA encoding NuABP and other genes of interest can be analyzed by northern analysis or microarray techniques.

XII. Production of NuABP Specific Antibodies

25 -

30

35

NuABP substantially purified using polyacrylamide gel electrophoresis (PAGE; see, e.g., Harrington, M.G. (1990) Methods Enzymol. 182:488-495), or other purification techniques, is used to immunize rabbits and to produce antibodies using standard protocols.

Alternatively, the NuABP amino acid sequence is analyzed using LASERGENE software (DNASTAR) to determine regions of high immunogenicity, and a corresponding oligopeptide is synthesized and used to raise antibodies by means known to those of skill in the art. Methods for selection of appropriate epitopes, such as those near the C-terminus or in hydrophilic regions are well described in the art. (See, e.g., Ausubel, 1995, supra. ch. 11.)

Typically, oligopeptides of about 15 residues in length are synthesized using an ABI 431A peptide synthesizer (Perkin-Elmer) using fmoc-chemistry and coupled to KLH (Sigma-Aldrich, St. Louis MO) by reaction with N-maleimidobenzoyl-N-hydroxysuccinimide ester (MBS) to increase

immunogenicity. (See. e.g., Ausubel, 1995, <u>supra.</u>) Rabbits are immunized with the oligopeptide-KLH complex in complete Freund's adjuvant. Resulting antisera are tested for antipeptide and anti-NuABP activity by, for example, binding the peptide or NuABP to a substrate, blocking with 1% BSA, reacting with rabbit antisera, washing, and reacting with radio-iodinated goat anti-rabbit IgG.

XIII. Purification of Naturally Occurring NuABP Using Specific Antibodies

Naturally occurring or recombinant NuABP is substantially purified by immunoaffinity chromatography using antibodies specific for NuABP. An immunoaffinity column is constructed by covalently coupling anti-NuABP antibody to an activated chromatographic resin, such as CNBr-activated SEPHAROSE (Amersham Pharmacia Biotech). After the coupling, the resin is blocked and washed according to the manufacturer's instructions.

Media containing NuABP are passed over the immunoaffinity column, and the column is washed under conditions that allow the preferential absorbance of NuABP (e.g., high ionic strength buffers in the presence of detergent). The column is eluted under conditions that disrupt antibody/NuABP binding (e.g., a buffer of pH 2 to pH 3, or a high concentration of a chaotrope, such as urea or thiocyanate ion), and NuABP is collected.

XIV. Identification of Molecules Which Interact with NuABP

10

20

NuABP, or biologically active fragments thereof, are labeled with ¹²⁵! Bolton-Hunter reagent. (See, e.g., Bolton A.E. and W.M. Hunter (1973) Biochem. J. 133:529-539.) Candidate molecules previously arrayed in the wells of a multi-well plate are incubated with the labeled NuABP, washed, and any wells with labeled NuABP complex are assayed. Data obtained using different concentrations of NuABP are used to calculate values for the number, affinity, and association of NuABP with the candidate molecules.

Various modifications and variations of the described methods and systems of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention. Although the invention has been described in connection with certain embodiments, it should be understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the described modes for carrying out the invention which are obvious to those skilled in molecular biology or related fields are intended to be within the scope of the following claims.

FABLE

33 SPLNFETO1 02 SYNORABO1 08 KIDNNOTO1 61 PLACNOBO1 61 HNT2RATO1 72 PROSNOTO2 00 COLNTUTO2 67 KIDNTUTO1 67 KIDNTUTO1 68 THYRNOTO3	Nucleotide Clone Library Fragments Fragments
2 57 079702 SYNORABOL 4 59 116208 KIDNROTOL 5 60 259161 HNTZRATOL 6 61 320087 BOSIHETOZ 7 62 491271 HNTZRATOL 8 63 585172 PROSNOTOZ 9 64 615200 COLNITUTOZ 10 65 997067 KIDNITUTOL 11 66 144326 HYRRANOTOZ	025733 SPLNFET01
3 58 116208 KIDNROTOL 4 59 179261 PLACNOBOL 5 60 259161 HNTZRATOL 6 61 320087 BOSIHETOZ 7 62 491271 HNTZAGTOL 8 63 585172 PROSNOTOZ 9 64 615200 COLNITUTOZ 10 65 997067 KIDNITUTOL 11 66 144326 HYRROTOL	079702
4 59 179261 PLACNOBOL	116208 KIDNNOT01
5 60 259161 HNT2RAT01 6 61 320087 EOSIHETO2 7 62 491271 HNT2AGT01 8 63 585172 PROSNOTO2 9 64 615200 COLNTUTO2 10 65 997067 KIDNTUTO1 11 66 144326 THYRNOTO3	179261
6 61 320087 EOSIHETO2 7 62 491271 HNT2AGTO1 8 63 585172 PROSNOTO2 9 64 615200 COLNTUTO2 10 65 997067 KIDNITUTO1 11 66 144326 THYRNOTO3	259161 HNT2RAT01
62 491271 HNT2AGT01 63 585172 PROSNOT02 64 615200 COLNTUT02 65 997067 KIDNTUT01 66 144326 THYRNOT03	320087
64 615200 COLNTUTO2 65 997067 KIDNTUTO1 65 144326 THYRNOTO3	491271 HNT2AGT01
64 615200 COLNTUTO2 65 997067 KIDNTUTO1 66 144326 THYRNOTO3	585172 PROSNOT02
65 997067 KIDNTUT01 66 144326 THYRNOT03	615200 COLNTUTO2
66 144326 THYRNOT03	125981X3 (1448201H1 1918706H1 2900479H1
010011112	144326 THYRNOTO3 1257005F1 (MENITUTO3), 1443262H1 2 2474133T6 (THPINOTO3), 3594075H1

	Protein SEQ ID NO:	Nucleotide SEQ ID NO:	Clone	Library	Fragments
	12	67	152164 8	BLADTUT04	292173H1 (TMLRJDT01), 819925R6 (KERANOTO2), 1353913H1 (LATRTUT02), 252168HH and 1527304FT (LADPUT04), 1961178H1 (RASTNOTO4), 234250gFT (TESTUTOTO), 4899370H1 (ONARDITO1), 504358FR2 (PLARTEREI)
	13	89	168549	PROSNOT15	903980X13, 903980X14 and 903980X17 (COLNNOTO7), 1685494H1 (PROSNOT15), 4164127T6 (BRSTNOT32)
	14	69	173082	BRSTTUT08	116146R1 (KIDNNOTG1), 836856R1 (PROSNOTG7), 1730829H1, 1730829X11C1, 1730829X12C1 and 1730829X12C1 (RRGYNOTG8), 1559889R6 (BRGYNOTG4), 188079H1 (PROSNOTG6), 3384625H1 (ESCGNOTG6)
	15	7.0	186464	PROSNOT19	1844972H1 (COLNNOTOB), 1864641F6 and 186464H1 (PROSNOT19), 3090702T6 (MSENDOT19), 3411665H1 (BRSTTUSOB), 5152366H1 (HRARERTOB), 5166179H1 (MUSCOWNOT)
	16	71	244460	THP1NOT03	1506658F1 (BRAITUTO7), 1532034F1 (SPLANNOTO4), 2444604H1 (THPINOTO3)
56	17	72	244500 8	THP1NOT03	605598712 (BRSTYTOI), 628644H (KIDNNOTOS), 73212481 (LUNGNOTO3), 81919481 (KERANOTO2), 129569H (HRBTYTOTA), 15632056 (LUNGNOTI2), 19031278 (BLADTYTOS), 2445008H (FHPINOTO3), 2681125H (SINIUTOTO)
	18	73	257246 2	HIPOAZT01	396323R6 (PITUNOTO2), 863622H1 (BRAITUTO3), 1848956F6 and 1848956F6 (LUNGEFRO3), 234584TH1 (FESTYUTO2), 2396384F6 (THP1AZPO1), 2572462H1 (HIPOAZPO1), 2650980F6 (LUNGTUT12), 2814325H1 (OVARNOT10), 5076051H1 (COLCTUTO2)
	19	74	257289 2	HIPOAZT01	030596XISR1 (THPINOBOI), 539564XII (INORNOTOZ), 1275514F1 and 125544F6 (TESTTUTOZ), 2112383H1 (BRAITUTOZ), 2572892H1 (HIPOAZTOI), 2966518H1 (CARGDITOI)
	20	75	278567	BRSTNOT13	261399H1 (HNT2AGT01), 1274739F1 (TESTTUT02), 2785674H1 (ERSTNOT13)
1	21	76	279747	NPOLNOT01	302614X13 (TESTNOTO4), 2797479H1 (NPOLNOTO1), SAIA02597F1, SAIA00739F1, SAIA00739F1,
	22	77	296064 0	ADRENOT09	027211R1, 027211X1 and 027211X3 (SPLNFBT01), 1401538F6 (BRAITUT08), 2496984F6 (ADRETUT05), 2960640H1 (ADRENOT09), 3211036T6 (BLADNOT08)
	23	78	345405	SPLNNOT11	279331R6 (LIVRTUTO2), 2515972R6 and 2516010T6 (LIVRTUTO4), 2910726F6 (KIDNTUT15), 3454051H1 (SPLNNOT11)

l					
щ ~-	Protein SEQ ID NO:	Nucleotide SEQ ID NO:	Clone	Library	Fragments
	24	79	351064 0	CONCNOT01	556354H1 (MPHGLPT02), 581085H1 (BRAVTXT05), 990636E6 (COLNNOTI1), 179115SF6 and 179918F6 and 179918FF6 (COLNNOT2), 3510640H1 (CONCNOT01), 4326648H1 (TYKNUND1), SSHAQ1099F1
	25	80	381508 3	TONSNOT03	2026951R6 and 2026951r6 (KERANOTO2), 2300211R6 (BR9TNOTOS), 2505283F6 (CONUTUPU), 3187267R6 (THYMNONO4), 3815083H1 and 3915083T6 (TONSKOTO3)
	26	81	398845 7	LUNGNON03	609622R6 (COLNNOTO1), 1710465F6 (PROSNOT16), 3988457H1 (LUNGNON03), SAQAO0089F1, SAQAQ3055F1, SAQRONO79F1
	27	82	131890	BMARNOT 02	131890H1 (BMARNOTO2), 131890T6 (BMARNOTO2), 132849R6 (BMARNOTO2), 3357071F6 (PROSTUTI6)
	28	83	238642	SINTNOT02	238642H1 (SINTNOFO2), 1620593F6 (BRAITUT13), 1620593H1 (BRAITUT13), 1620593T6 (BRAITUT13), 2534087F6 (BRAINNT18)
5	29	84	669862	CRBLNOT01	347231X7 (THYMNOTO2), 66962H1 (CRBLNOTO1), 2224458R6 (HIPONONO2), 4224448F6 (HIPONONOX), 2622610T6 (KERANOTO2), 3536262H1 (KIDNNOT25), 4204212H1 (REALTOT25)
7	30	85	100366 3	BRSTNOT03	850478T1 (NGANNOT01), 1003663H1 (BRSINOT03), 1252179F2 (LUNGFET03), 1293336F1 (PGANNOT03), 1813002F6 (PDESEMBL)
	31	86	143255	BEPINON01	1432557H1 2182184F6
	32	87	144177 0	THYRNOT03	HUVENOBO1), 1441770H1 (UTRSNOT11), 4533672H1
	33	88	145668 4	COLNFET02	
	34	68	160291 6	BLADNOT03	3397976X305D2 (UTRSNOT16)
	35	06	169281 6	COLNNOT23	999017R6 (KIDNTUT01), 1342490T1 (COLNTUT03), 1421981F1 (KIDNNOT09), 1692816H1 (COLNNOT23), 2176832F6 (FRIPTUROFOR) 3451404FE (FRIPTUROFOR)
	36	91	196819 1	BRSTNOT04	
\Box	37	92	205206	LIVRFET02	3931936F6

	Protein SEQ ID NO:	Nucleotide SEQ ID NO:	Clone	Library	Fragments
	38	93	205620	BEPINOT01	071525F1 (PLACNOBOL), 162720R1 (ADENINBOL), 270408H1 (HATZNOTO), 1477837H1 (CORNOVOZO), 1931058F6 (COLNIVUTO3), 2056207H1 (BEFINOTOL), 2056207X11AL (BREFINOTOS), 2231066F6 (PROSNOTIS), 2420063X309D4 (SCORNONOZ), 3424630H1 (BRSTNOROL), 3873760F6 (HEARNOTOS), 3873760F6 (HEARNOTOS), SBOAGO627D1,
	39	94	210180 3	BRAITUT02	399584R6 (PITUNOTO2), 399584T6 (PITUNOTO2), 1649058F6 (PROSTUTO9), 1902809F6 (OVANDOTO), 2101803H1 (BRAITUTO2), 2101803R6 (BRAITUTO2), 2009653H1 (CRENATOTO)
	40	98	211236	BRAITUT03	946628R1 (PANCNOTOS), 1209447T1 (BRSTNOTO2), 1814624F6 (PROSNOT2)), 1113.824H (BRATUTUTO3), 29.82665H1 (BRALUPUTO3), 29.82665H1 (BRALOMOTOS), 556403H1 (ESOGTWOID), 5032729H (ENDINNOT), 60447H1 (ENDINNOTO)
I	41	96	211734	BRST-TUT02	2458342F6 (
8	42	97	211991	BRSTTUT02	2791421F6 (COLNTUT16),
	43	86	212345 6	BRSTNOT07	484031H1 (HWTZRATO1), 617559F1 (PGANNOTO1), 57559A1 (PGANNOTO1), 15759771 (LMONOTO3), 298712H1 (ARRNOTO1), 376496LH1 (RRSTWOTO7),
	44	66	214879	BRAINOT09	
	45	100	275194 3	THP1AZS08	1720187X16C1 (BLADNOT06), 2751943H1 (THP1AZSO8), 3492378H1 (ADRETUTO7)
	46	101	312891 3	LUNGTUT12	2551859F6 (LUNGTUT06), 3128913H1 (LUNGTUT12), SBMA01861F1, SBMA02298F1, SBMA01013F1, SBMA02403F1, SBMA01362F1
	47	102	328294	HEAONOT05	154741R6 (THPIPLBO2), 155904R6 (THPIPLBO2), 15781R1 (THPIPLBO2), 97920H1 (TONGOTUTO1), 1233933FG (LUNGFETO3), 165707FG (URETYUTO1), 2445017FG (THPINOTO3), 3282941H1 (HEAONOTUS), 3341633H1 (PELNNOTUS), 3517140H1 (LUNGRONOT)
	48	103	328665 6	HEAONOT05	898123H1 (BRSTWOT05), 3286656H1 (HEAONOT05), 3641429F6 (LUNGNOT30), 3657668F6 (ENDPNOT02)

Protein SEQ ID NO:	Nucleotide Clone SEQ ID NO: ID	Clone	Library	Fragments
49	104	349080	EPIGNOT01	2238441H1 (PANCTUTO2), 2700133F6 (OVARTUTIO), 2700133F6 (OVARTUTIO), 3490226H1 (EPIGNOTO1), 3490802H1 (PROSTUTI)
50	105	350736 6	CONCNOT01	2130284H1 (KIDNNOTOS), 3507366H1 (CONCNOTO1), 3557087F6 (LUNGNOT31), 4241774H1 (SYNWDIT01)
51	105	357306 0	BRONNOT01	3573060F6 (BRONNOT01), 3573060H1 (BRONNOT01), 3573060T6 (BRONNOT01), 3867263H1 (BRAITUT07), 5013346H1 (BRAXNOT03)
52	107	357366 1	BRONNOT01	3028034F6 (HEARFET02), 3152642H1 (ADRENON04), 3573661F6 (BRONNOT01), 3573661H1 (BRONNOT01), 3577568F6 (BRONNOT01)
53	108	363342 2	LIVRNOT03	033412B6 (THELNOBOL), 074123F1 (THELPEBOL), 263241H1 (HUNZAGTOL), 7485FR1 (BASITTUROL), 129208BT1 (PGANNOTO3), 1517449T1 (PANNOTOGL), 363342BH1 (LIVUNOTO3)
54	109	399337 7	TUNGNON03	3003233H1 (TLYMNOT06), 3993377H1 (LUNGNON03), 3993377T6 (LUNGNON03), 4251662F6 (BRADDIR01), SBSA02001V1
55	110	471793 6	BRAIHCT02	4717936F6 (BRAIHCT02), 4717936H1 (BRAIHCT02), 4717936T6 (BRAIHCT02)

tide Acid Phosphorylation Olycosylation Signature Sequence(s) Homologous Sequence Se							
No. Residues 1168 Saguence(s)	Folype tide	Ω.	Potential Phosphorylation	Potential Glycosylation	Signature	Identification/	Analytical
2 593 870 860 784 11 754 7134 8308 7132 2 593 7136 8313 2 593 7136 8314 3 534 838 843 840 3 534 843 843 840 4 255 7136 841 831 5 61 738 744 833 6 738 746 842 7 738 746 842 7 738 748 843 840 7 738 748 843 840 7 738 748 843 840 7 738 748 843 840 7 738 843 843 840 7 738 843 843 840 7 748 843 843 840 7 757 748 841 841 7 758 718 841 841 7 758 718 841 841 7 758 718 718 718 718 7 748 748 748 748 748 7 758 748 748 748 748 7 758 748 748 748 748 7 758 748 748 748 748 7 758 748 748 748 748 748 7 758 748 748 748 748 748 748 748 748 748 74	NO:		Sires	Sites	Sequence(s)	Homologous Sequence	Databases
1 754 F12 S10. 814 112 S11. 8146 F16 F18 T1846 F16 F18 T1846 F16 F18 T1846 F16 F18 T189 F1846 F17 F18 F18 T189 F18 F18 F18 F18 F18 F18 F18 F18 F18 F18			S6 T15 S22 S26				
1 754 113 5121 5146 1 754 7136 7138 7184 1 754 7136 7139 7134 1 754 7136 7139 7134 1 754 7136 7139 7134 2 593 713 571 2 593 713 571 2 713 572 51 3 534 545 545 545 3 534 545 545 4 255 714 68 716 1 754 716 7174 3 534 545 545 4 255 714 68 716 1 755 714 7159 716 2 745 715 716 3 746 716 717 4 255 714 718 715 5 714 718 718 718 7 748 718 718 719 7 752 719 719 7 752 719 719 7 752 719 719 7 752 719 719 7 752 719 719 7 752 719 719 7 752 719			528 530 S60 Y84				
11.2 \$1.2 \$1.2 \$1.4 \$1.5 \$1.5 \$1.5 \$1.5 \$1.5 \$1.5 \$1.5 \$1.5			238 STOZ STO3				
1 754 7126 71284 1134 1 754 7136 71284 71284 1255 5303 1 754 7136 2139 71394 PFAM 7136 7139 7139 71394 PFAM 2 593 713 5751 2 593 713 575 713 7140 542 542 528 715 575 713 575 713 715 575 713 7140 7140 717 719 719 719 719 717 719 719 719 719 718 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719 719			S112 S121 S146				
1 754 7304 7305 73.7			T166 T183 T184				
1 754 7304 3308 7327 PPAM 7309 7448 8496 7508 7448 8496 7508 8615 867 8608 8615 867 8619 8711 8711 8711 8711 8711 8711 8711 87			Y253		2000		PFAM.
Tight Tigh	1	754	T304 S308 T327		1435 - P474	DNA polymerase	BLOCKS
1399 T448 5496 1399 T448 5496 1307 1448 5496 1307 5597 5672 5673 1318 5731 1318 5734 131		_	T361 T393 T394		PFAM		MOTTES,
Y58 S56 F608			T399 T448 S496				
2 593 713 8751 2 185 862 8673 3 2 182 713 8751 3 2 182 713 8751 3 11 1159 7160 3 185 224 8288 3 1718 834 8333 4 61 479 7483 3 738 838 8479 3 534 8479 848 3 7488 756 3 748 8479 846 3 534 848 8479 846 4 255 7248 7168 7169 7 725			Y583 S586 T608				
S682 S691 S711 S713 S721 T65 S83 S12 V62 T65 S83 S14 T152 T160 S165 S693 T160 S165 S294 S286 T311 S314 S333 T416 S308 S401 T410 S426 S452 T527 T480 S490 T527 T548 T68 Y169 T547 T68 Y169 T548 Y169 Y169 T548 Y169 Y169 T548 Y169 Y169 Y169 T548	_		S635 S672 S673				
S713 S751 S714 S751 S717 S751 S717 S716 S716 S716 S716 S716 S716 S716			S682 S691 S711				
2 593 T1 1159 T160 2 186 5245 5288 2 186 5245 5288 2 186 5245 5288 2 1871 5314 5333 2 1871 5314 5333 2 1871 5314 5333 2 1871 5314 5333 2 1871 5314 5314 2 534 5419 5425 2 1871 5314 5314 2 534 5419 5410 2 1871 5314 5419 5410 2 1871 5314 5419 5410 2 1871 5314 5419 5410 2 1871 5314 5419 5410 2 1871 5314 5419 5410 2 1871 5314 5419 5410 2 1871 5314 5419 5410 2 1871 5314 5419 5410 2 1871 5314 5419 5410 2 1871 5410 5410 2 1871 5410 5410 2 1871 5410 5410 2 1871 5410 5410 2 1871 5410 5410 2 1871 5410 2			S713 S751				
2 593 T311 5314 533 2 593 T311 5314 533 T316 5360 5401 T40 5426 5452 5461 5479 T483 T40 525 531 T314 T39 522 5301 T344 T39 522 5301 T346 T39 540 540 FFAM T527 T460 5490 T527 T460 5490 T527 T460 5490 T527 T460 5490 T527 T340 T168 Y169 T169 T141 - H163 T249 T168 Y169 T168 Y169 T169 - H219 T255 T240 T255 T240 T264 T264			S32 Y62 T65 S83				
1185 254 2588 E102 - A157 1716 314 5333 E202 - A157 1716 3280 5401 C252 1716 328 5452 C201 - C252 1717 32 525 3301 7344 C201 - C252 1717 - D57 334 3517 340 5425 C201 - C252 1717 - D57 334 3517 340 5425 C201 - C252 1718 525 331 7404 5425 C252 3117 - D57 311	50		S141 T159 T160				
171 1814 8333 171 1814 8348 8348 8461 8479 4883 8461 8479 4883 8461 8479 4883 8461 8479 8483 8479 4883 8478 84			S185 S254 S288				
7176 8380 8401 4201 - C252 3461 8477 748 7176 841 8425 534 8438 8440 PFAM 71527 7480 8490 PFAM 71527 7480 7480 PFAM 71527 7480 PFAM 71527 PFAM	2	593	T311 S314 S333		E102 - A15/	PHD finger DNA binding	BLAST DEAM
740 822 6352 3461 3479 7483 7485 7576 719 582 8301 7344 534 8418 8439 840 7527 7527 7527 7527 7527 7527 7527 752		3	T368 S380 S401		Q201 - C252	protein (GI 3342452)	MOTTES
1465 15479 17483 1465 15461 1844 179 552 8301 17344 179 552 8301 17344 179 552 8301 17344 179 552 8303 17404 6425 170 6426 170			T410 S426 S452			77	211100
T485 T786 T485 T787 T79 S52 S301 T344 T79 S52 S301 T344 T79 S52 S301 T344 T79 T404 S425 T77 T408 S430 T75 T408 S430 T75 T408 S430 T75 T408 S430 T75 T408 T168 T169 T169 T169 T169 T169 T169 T169 T169			S461 S479 T483				
179 552 530 T 344 534 5438 5439 5445 5438 5439 5440 5438 5439 5440 FFAM F			T485 T576				
\$34 \$4404 \$425 \$117 - D57 \$473 \$7400 \$4405 \$470 \$7405 \$440 \$440 \$8473 \$7480 \$490 \$7527 \$758 - C86 \$149 T168 Y169 \$759 \$759 - H191 \$759 \$759 - H191 \$759 \$750 \$750 \$750 \$750 \$750 \$750 \$750 \$750			T39 S52 S301 T344				
534 5439 5440 11/ - D5/ 5473 7480 5490 FFAM 7527 7480 5490 FFAM 7527 7480 7168 7169 7169 - H191 7248 7168 7169 7169 - H219 7249 7168 7169 7169 - H219 7255 7225 - H247			S373 T404 S425				DFAM
S473 T480 S490 PFAM T527	3	534	S438 S439 S440		11/ - D5/	DNA polymerase	BIOCKS
255 T248 T189 T189 T189 - R180 T189 T189 T191 T191 T191 T191 T191 T191			S473 T480 S490		PFAM	0.000	MONTES,
255 T248 T168 Y169 F161 Y169 - C86 F141 - H163 F141 H163 F141 - H163 F141 - H163 F141 - H163 F141 - H163 F141 F141 F141 F141 F141 F141 F141 F14			T527				FOLLES
255 T248 T168 Y169 F141 - H163 Y169 - H21 - H219 Y197 - H219 Y225 + H219 Y225 - H247 Y225 - H247			S10 S26 Y35 Y113		Y58 - C86		
255 T248 X169 - H191 Y199 - H219 Y225 - H247 Y225 - H247			S149 T168 Y169		F141 - H163		BLAST. PFAM.
1197 - H219 1225 - H247 1255 - H247	4	255	T248		Y169 - H191	CZHZ-type zinc finger	BLOCKS
H247	_				Y197 - H219	procein	PRINTS.
PPAM					Y225 - H247	(GI 498721)	MOTIFS
					PFAM		

PCT/US00/02237

			(2002) = 777777	(2000)		
Polypep tide Seq ID NO:	Amino Acid Residues	Potential Phosphorylation Sites	Potential Glycosylation Sites	Signature Sequence(s)	Identification/ Homologous Sequence	Analytical Methods and Databases
Ln.	562	T29 T43 S76 S142 S165 S202 T214 Y302 S305 Y349 S385 S500 T526 S527		Y235 - D312 PFAM	DNA helicase (GI 531243; SEQ ID NO:113)	BLAST, PFAM, BLOCKS, MOTIFS
v	432	S33 S58 T166 T172 S197 T230 T261 S275 S286 S290 S298 S338 T362 S376 T407 T409		E329 - A355 BLOCKS	CCAAT-box-binding transcription factor	BLOCKS, MOTIFS
7	799	T24 S33 T43 S73 T88 S91 S110 S147 T219 S262 S323 Y380 S532 S586 S756 T795		H250 - H291 Y324 - H346 Y253 - H374 Y380 - H402 PPAM	C2H2-type zinc finger protein (GI 498727)	BLAST, PFAM, BLOCKS, PRINTS, MOTIFS
80	137	S3 T38 S74 S75 S118		R85 - L97 PFAM	BTB domain/C2H2-type zinc finger protein	PFAM, PRINTS, MOTIFS
6	230	T178 S187			sirtuin type 3 (GI 5225322)	BLAST, MOTIFS
10	446	T3 S28 S32 T52 T94 T96 S135 S143 T159 T165 S171 S433		H200 - H222 Y228 - H250 Y256 - H278 Y284 - H306	zinc finger protein ZFP113 (GI 5640017)	BLAST, PFAM, BLOCKS, PRINTS, MOTIFS
11	428	\$72 \$92 \$101 \$118 \$120 \$125 \$7245 \$127 \$289 \$315 \$317 \$726 \$409			Skeletal muscle BOP2 (GI 5870834; SEQ ID NO:117)	BLAST, MOTIFS

			IABLE 2 (cont.)	(cont.)		
Polypep tide Seg ID NO:	Amino Acid Residues	Potential Phosphorylation Sites	Potential Glycosylation Sites	Signature Sequence(s)	Identification/ Homologous Sequence	Analytical Methods and Databases
12	290	S45 Y52 T60 S83 S90 T95 T116 T145 T233 T330 S391 S410 S411 T420 T439 T490 S521			Methyl-CpG binding protein (GI 2239126)	BLAST, MOTIFS
13	479	515 Y29 S30 S118 T173 T183 S203 S217 Y232 S235 T255 S352 S362 Y451		Y232 - H254 H283 - H305 Y311 - H333 Y339 - H361 PFAM	SRE-ZBP (GI 936603)	BLAST, PFAM, BLOCKS, PRINTS, MOTIFS
14	433	S92 S96 T250 S319 T322 T327 S335 T344		C380 - C421 PFAM	C3HC4-type zinc finger protein	PFAM, BLOCKS, MOTTES
15	320	T6 S27 T125 T172 S229 S232 T239 S248 S259 S266 Y267 S291			Zinc finger factor (GI 3150148)	BLAST, MOTIFS
16	179	S11 T21 S46 S140			Single-stranded DNA binding protein (csdp)	BLAST, MOTIFS
17	494	T73 Y80 S104 Y116 T192 S289 S297 T329 T364 T376 S387		C13 - H41 BLOCKS	Zinc finger transcription factor (GI 2895870)	BLAST, BLOCKS, MOTIFS
18	401	S4 S82 S97 T166 S188 S249 S279 S299 S299 S294 S319 S368 S371 S372 S378 T392 S396		PS - K81 A114 - S179 G186 - P262 PFAM	HP1-BP74 (GI 1480112)	PFAM, BLOCKS, PRINTS, MOTIFS
						-

			I ADLE 2 (COIII.)	(courts)		
Polypep tide Seg ID NO:	Amino Acid Residues	Potential Phosphorylation Sites	Potential Glycosylation Sites	Signature Sequence(s)	Identification/ Homologous Sequence	Analytical Methods and Databases
19	264	S11 S25 S76 S82 S90 S92 S96 S119 T229		F154 - H176 C180 - H202 F208 - H230 Y236 - C259 PPAM	C2H2-type zinc finger protein (GI 429188)	BLAST, PFAM, BLOCKS, PRINTS, MOTIFS
20	153	T23 S40 T44 S110 S120 T124		R42 - E141 PFAM	High mobility group- like nuclear protein (GI 2822179)	BLAST, PFAM, BLOCKS, PRINTS, MOTIFS
21	243	S20 S21 S76 S100 S104 S160 T194 S196 S212 T222 Y229	÷	Y90 - H112 H118 - H140 Y146 - H168 Y174 - H195 PFAM	C2H2-type zinc finger protein (GI 38015)	BLAST, PFAM, BLOCKS, PRINTS, MOTIFS
22	485	T29 S34 S104 S147 T162 T248 S249 S256 S347 S452 S477		S309 - H331 H337 - H359 Y365 - H387 Y393 - H415 PFAM	BTB domain/C2H2-type zinc finger protein (GI 2843171)	BLAST, PFAM, BLOCKS, PRINTS, MOTIFS
23	160	S118		C5 - F62 C80 - F137 PFAM	LIM domain protein/CRP2 (GI 487284)	BLAST, PFAM, BLOCKS, PROFILESCAN, MOTIFS
24	511	S10 T36 S75 S90 S222 T245 T259 S399 S405 Y443 S500	-a	Y171 - P223 Y267 - K294 BLOCKS	2'-5'oligoadenylate synthetase-related protein p56 (GI 4731857)	BLOCKS, MOTIFS
25	310	S24 S39 T69 Y104 S185 T282 T296	-		SIR2 family transcriptional regulatory protein (GI 2648874)	BLAST, MOTIFS

TABLE 2 (cont.)

	Polypep tide Seq ID NO:	Amino Acid Residues	Potential Phosphorylation Sites	Potential Glycosylation Sites	Signature Sequence(s)	Identification/ Homologous Sequence	Analytical Methods and Databases
,,,	26	331	T29 S40 S74 S257 T270 S301		V112 - R134 BLOCKS	Histone protein	BLOCKS, MOTIFS
"	27	200	T43 S123 T129 S167 S183 S184			Zinc finger protein (g1373394)	MOTIFS
	28	100	S44 Y25 Y98		Transcription anti-terminator; bglG family:	transcription elongation factor (94336506)	MOTIFS BLAST BLOCKS
L.			S204 T487 S29 S34	N24 N52 N100	C2H2 zinc		
			T48 T227 S327	N481	fingers:		
			T367 T423 S483		Y191-H213		
			Y39 Y44 Y112 Y163		Y247-H269		MOTIFS
64	29	528			Y275-H297	Zinc finger protein	BLAST
					Y331-H353	(g498721)	BLOCKS
					Y387-H409		PFAM
					Y415-H437		
					Y443-H465		
_					Y471-H493		
_			T264 S305	N33 N79			MOTIFS
	0.5	Cuc			C3HC4 RING	C3HC4/BING zinc finger	BLAST
<u>'</u>	2	200			finger: C230-C271	protein (g1321818)	BLOCKS
							PROPTLECON
			S51 T94 S121 S123				
~	31	315	S142 S143 T184			Similar to	MOTIFS
			S232 S252 T36 T46			protein (#1947129)	BLAST
			S159 S163 S168			Process (9124/129)	
			S36 S56 T93 S104	N98 N103	4IZq		OUT BOX
<u></u>	32	120			transcription		DEAM
					factor:	-	BLOCKS
j					201 171		

	Analytical Methods and Databases	MOTIFS BLAST PFAM BLOCKS	MOTIFS BLAST BLOCKS PFAM PROFILESCAN	MOTIFS BLAST PFAM	MOTIFS BLAST PFAM	MOTIFS BLAST BLOCKS PFAM	MOTIFS BLAST BLOCKS PFAM	MOTIFS BLAST
	Identification/ Homologous Sequence	Zinc finger protein (g220643)	CHOX M product; homeobox protein (g62701)	geminin (g3219357)	Smarcel-related protein (GI 4321968)	BKLF; CACCC-box binding protein (g1244515)	ARI (RING finger) protein (g2058299)	skm-BOP2 zinc finger protein (g1809327)
(contro)	Signature Sequence(s)	C2H2 zinc fingers: C143-C171 Y169-H191 F197-H219	Homeobox: R14-K70	bZIP transcription factor: K115-E140	HMG box: M1-Q36	C2H2 zinc fingers: F278-C306 Y304-H328 F334-H356	C3HC4 RING fingers: C74-P120 N228-C235	
(control of (control	Potential Glycosylation Sites	N209		N18	N92	N45 N340	N337 N374 N388	N169 N206
	Potential Phosphorylation Sites	S59 S38 T207 S284 T319 T43 S80 T137 T155 T211 S238 T239	S80 T89 T39 T53	S176 T180 S184 T193 S201 S4 T25 S49	S79 S107 T127 T202 S45 S56 S124 T152 Y35	T329 T50 S125 S224 S230 S235 S344 S31 S215 S312 Y42	S68 T87 S153 S339 N337 N374 N388 S405 S55 T105 S315 S422 Y419	S283 T44 T57 T123 S136 T185 T220 S239 T268 S313 S330 T105 T109 S125 T216
	Amino Acid Residues	326	106	209	212	359	445	433
	Polypep tide Seq ID NO:	33	34	<u>د</u> د	36	37	38	39

				()		
Polypep tide Seg ID NO:	Amino Acid Residues	Potential Phosphorylation Sites	Potential Glycosylation Sites	Signature Sequence(s)	Identification/ Homologous Sequence	Analytical Methods and Databases
40	355	T72 T84 T184 S191 T244 S88 T162 T229 T294 S330	N308 N324		Sir2 family protein (g5353746)	MOTIFS BLAST
41	443	T76 T106 S148 T309 S12 T253 S299 S357 T373 T427 Y61 Y114	N366 N425	Myb-like DNA binding domain: D72-F118	ADA2 transcriptional adaptor protein (g170991)	MOTIFS BLAST BLOCKS PFAM PROFILESCAN
42	164	S37 T55 T64 S83 S22 Y146	N117	mutT domain: V29-L70		MOTIFS BLOCKS PFAM
43	215	S79 S127 T12 T45		HMG box: M1-Q36	Sry-related protein (g211510)	MOTIFS BLAST PFAM
4	539	T15 524 566 583 197 T105 T109 5128 T149 5153 5188 T203 T225 T238 T296 5466 Y258	N465	C2H2 zinc fingers: fingers: Y256-H280 Y256-H208 Y342-H306 Y342-H306 Y396-H420 Y425-H604 Y425-H504	C2H2 zinc finger protein (95757625)	MOTIFS BLAST PFAM BLOCKS
45	182	T59 S112 S120 S100 S139 Y64			Transcriptional regulator (g2621798)	MOTIFS BLAST
						The same of the sa

ŭ	Polypep	Amino	Potential	Potential			Acces 10.00 (10.00)
T.	tide	Arid	Phosphorylation	Glycosylation	Signature	Identification/	Marytical
χ Σ	Seq ID NO:	Residues	Sites	Sites	Seguence(s)	Homologous Sequence	Methods and Databases
_			S494 S31 S44 S117		C2H2 zinc		
_			T123 S185 S216	N351	fingers:		
			S476 S504 S176		Y285-H307		
			S182 S211 T249		Y313-H335		07.4
			S293 S323 S409		Y341-H363		MOLIES
46	·	534	T489 Y76 Y285		C369-H391	And tinger process	BLAST
					Y397-H419	(813/3384)	Pram
_					X425-H447		BEOCEAS
					Y453-H475		
					Y481-H503		
					Y509-H531		
			S5 S7 S40 S45 S46 N	N44 N177			MOTIFS
			S100 S144 S26		Myc-type HLH		BLAST
47	7	206	S107 T148 S185		domain:	Musculin (g3599519)	PFAM
7			Y38		Q108-R160		BLOCKS
_							PROFILESCAN
48	•	172	T5 S87 S96 S115			KRAB zinc finger	MOTIFS
1			T124 S22 T64			protein (g1049295)	BLAST
			S185 S14 S48 T54	N210 N214 N238	C2H2 zinc		MORTEO
_ :			S118 T139 T161	N260	fingers:	Repressor	E POT TO
49	•	275	T189 T217 Y256		F172-H194	transcriptional factor	DUASI
					Y200-H222	(g1017722)	FIRM
_					Y228-H250		BLOCKS
			S157 S42 T167	N40	C3HC4 RING		MOTIFS
20	_	236	T222 T81 Y213		finger:	Arlagne-2 KING tinger	BLAST
					P126-L150	procein (g3445441)	BLOCKS
			S7 S8 S116 T127	N2	herman	200	MOTIFS
21		214	S154 S191 T31 S41		V113-E134	(d833629)	BLAST
			1204				BLOCKS

Polypep Amino Potential Potential Potential Lider Potential Pote						, , , , ,		
Phosphorylation Glycosylation Signature Identification Seq ID Residues Sites Sites Sites Sites Homologous Sequence So		Polypep	amin's	Potential	Potential			
Seq ID Residues Sites Sequence (s) Homologous Sequence NO: 139 T201 S118 N2 C3HC4 RING Sequence Sequence Si26 S26 T313 T35 S315 S39 T35 S315 S39 T35 S315 S39 S316 S39 T441 S316 S316 S39 T441 S316 S316 S39 T441 S316 S316 S39 T441 S316 S316 S39 T442 Y48 S316 S316 S39 S412 T42 Y48 S316 S316 S316 S316 S316 S316 S316 S316		tide	Acid	Phosphorylation	Glycosylation	Signature	Identification/	Analytical
134 1392 5118 N2		Seq ID	Residues	Sites	Sites	Sequence(s)	Homologous Sequence	Databases
1784 1792 518 NZ 1794 RING 1794 RING 1794 RING 1795 1791 S18 NZ 1795 1791 S18 NZ 1795 1791 S18 NZ 1795 1791 S18 NZ 1795 1795 S175 S175 S175 S175 S175 S175 S175 S17		NO:						
119 720 1187 220 C3H64 RING Midline I I Cerebellar				T348 T392 S118	N2		100	MOTIFS
52 396 5204 580 5112 C26-C50 C26-C50 C26-C50 C26-C50 C26-C50 C26-C50 C26-C50 C26-C50 C26-C50 C320 520 5215 5213 C26-C50 C320 5215 5213 C26-C50 C320 5215 5213 C320 5215 5213 C320 5215 5213 C320 5213 5213 5215 5213 C320 5213 5213 5215 5213 5215 5213 5215 5213 5215 5213 5215 5215				T193 T201 S270		C3HC4 RING	Midilne I/ cerebellar	BLAST
S206 S260 T313 C26-C50 Parcelan 1355 S315 S389 C26-C50 Parcelan 1355 S315 S389 C378 S315 S339 C38-C50 C38-C50 C38-C50 1318 S412 T502 S68 N25 N66 N246 C48-C50 C4	_	52	396	S294 S80 S112		finger:	Isolorm I KING Ilnger	PFAM
1735 5387 1946 1728 1735 5387 1948c25033 1948c25033 1957 5387 1957 5389 1957 5329 1957 5329 1957 5329 1957 1739 1739 1741 1957 1	_			S206 S260 T313		C26-C50	protein	BLOCKS
53 486 729 718 5156 529 718 5146 5 - mucleotidase 1721 5425 5 - mucleotidase 1722 719 7286 1725 7441 1724 719 5148 1725 7441 1724 719 5148 1725 719 728				T355 S375 S387			(g3462503)	PROFILESCAN
53 486 T271 5425 5779 5146 5770				S29 T58 S155 S239				
T211 8425 1925 866 1925 866 1925 866 1925 866 1925 866 1925 866 1925 866 1925 866 1925 866 1925 862 1925 825 1927 7441 1925 825 1927 7441 1925 825 1927 825 1927 825 1927 825 1927 825 1927 825 1927 825 1927 825 825 825 825 825 825 825 825 825 825		53	486	T292 T379 S146			5'-nucleotidase	MOTIFS
54 555 T319 7127 7421 811 7257 7274 81 727 7274 8 812 727 7434 81 7257 7274 81 7274 7274 7274 81 7274 7274 7274 7274 7274 7274 7274 727				T271 S425			(9633071)	BLAST
S195 T199 T226 N364 ATP/GTP binding tremastripon factor I				S432 T502 S68	N25 N66 N246			
S45 S55 T34 T441 ATP/GTP binding Cermination factor I				S195 T199 T226	N364		Transcription	
54 555 T534 S170 S248 site (P-loop): (TTF-I) interacting 228 2591 T327 pp.tide 5 isoform 738 5391 542.				S315 T379 T441		ATP/GTP binding	termination factor I	
73.72		54	555	T534 S170 S248		site (P-loop):	(TTF-I) interacting	MOTIFS
T34 742 Y48				S282 S291 T327		A434-T441	peptide 5 isoform	BLAST
T481 Y257 Y274 Putative leucine-rich T34 T42 Y48 DNA-binding protein (455591)	(T336 S391 S422			(42183083)	
61 T34 T42 Y48 Putative leucine-rich DNA-binding protein (455591)	58			T481 Y257 Y274				
61 DNA-binding protein (q555991)				T34 T42 Y48			Putative leucine-rich	
_		22	61				DNA-binding protein	MOTIFS
	_						(g555991)	BLAST

		TABLE 3		
Nucleotide Seq ID NO:	Selected Fragment	Tissue Expression (Fraction of Total)	Disease or Condition (Fraction of Total)	Vector
56	169-215	Reproductive (0.224) Nervous (0.198) Cardiovascular (0.112)	Cell Proliferative (0.725) Inflammation (0.190)	PBLUESCRIPT
57	551-595	Hematopoietic/Immune (0.240) Reproductive (0.180) Gastrointestinal (0.120)	Cell Proliferative (0.700) Inflammation (0.360)	PBLUESCRIPT
58	541-585	Nervous (0.286) Reproductive (0.286) Cardiovascular (0.214)	Cell Proliferative (0.643) Trauma (0.214)	PBLUESCRIPT
59	109-153	Reproductive (1.000)	Cell Proliferative (1.000) Inflammation (1.000)	PBLUESCRIPT
09	435-479	Hematopoietic/Immune (0.211) Gastrointestinal (0.183) Reproductive (0.183)	Cell Proliferative (0.620) Inflammation (0.338)	PBLUESCRIPT
61	1195-1239	Reproductive (0.248) Cardiovascular (0.174) Nervous (0.157)	Cell Proliferative (0.637) Inflammation (0.256)	PBLUESCRIPT
62	217-261	Reproductive (0.429) Nervous (0.238) Cardiovascular (0.095)	Cell Proliferative (0.667) Inflammation (0.143) Trauma (0.095)	PBLUESCRIPT
63	919-963	Reproductive (0.265) Nervous (0.235) Cardiovascular (0.088)	Cell Proliferative (0.618) Inflammation (0.206)	PSP0RT1
64	823-876	Reproductive (0.382) Nervous (0.176) Gastrointestinal (0.118)	Cell Proliferative (0.794)	PSPORT1
65	380-424	Reproductive (0.346) Nervous (0.154) Gastrointestinal (0.135)	Cell Proliferative (0.750) Inflammation (0.231)	PSPORT1

TARLE 3 (conf.)

Mucleotide Selected Fragment Seq ID NO: 757-801		Tissue Expression	Disease or Condition	
			(Fraction of Total)	Vector
		Nervous (0.222)	Cell Proliferative	DINCY
		Reproductive (0.167)	Inflammation (0.222)	
	Nev	Reproductive (0.246)	Cell Proliferative	
	-	Nervous (0.180)	(0.639) Inflammation	pINCY
	Gas	Gastrointestinal (0.148)	(0.246)	
		Reproductive (0.500)	Cell Proliferative	
	Nez	Nervous (0.200)	(0.700)	pINCY
	Gas	Gastrointestinal (0.150)	Trauma (0.150)	
		Reproductive (0.278)	Cell Proliferative	
	Hen	Hematopoietic/Immune (0.204)	(0.777) Inflammation	DINCY
	Nez	Nervous (0.148)	(0.222)	
		Reproductive (0.261)	Cell Proliferative	
	Gas	Gastrointestinal (0.217)	(0.565)	PINCY
	Neı	Nervous (0.174)	Trauma (0.130)	
71	_	Nervous (0.250)	Cell Proliferative	
	Der	Developmental (0.208)	(0.583)	PINCY
	Gas	Gastrointestinal (0.167)	Trauma (0.167)	
529-573		Reproductive (0.186)	Cell Proliferative	
72	Gas	Gastrointestinal (0.168)	(0.700)	PINCY
	Her	Hematopoietic/Immune (0.138)	Inflammation (0.251)	
1784-1828		Reproductive (0.286)	Cell Proliferative	
73	Her	Hematopoietic/Immune (0.190)	(0.667) Inflammation	PSPORT1
	Ne	Nervous (0.167)	(0.286)	
111-155		Reproductive (0.316)	Cell Proliferative	
74	Ne		(0.632) Inflammation	PSPORT1
	Her	Hematopoietic/Immune (0.158)	(0.211)	
543-587		Reproductive (0.258)	Cell Proliferative	
75	Ne	Nervous (0.206)	(0.608) Inflammation	PINCY
	Gas	Gastrointestinal (0.134)	(0.196)	

	-	
	5	
,	٤	
	,	
1	Ġ	ì
	<	
ł		

		TABLE 3 (cont.)		
Mucleotide Seq ID NO:	Selected Fragment	Tissue Expression (Fraction of Total)	Disease or Condition (Fraction of Total)	Vector
76	272-316	Reproductive (0.246) Nervous (0.180) Hematopoietic/Immune (0.148)	Cell Proliferative (0.606) Inflammation (0.279)	pincy
77	227-271	Hematopoietic/Immune (0.222) Endocrine (0.167) Cardiovascular (0.111)	Cell Proliferative (0.666) Inflammation (0.333)	pINCY
78	487-531	Gastrointestinal (0.375) Reproductive (0.250)	Cell Proliferative (0.500) Inflammation (0.250)	pINCY
79	111-155	Gastrointestinal (0.280) Hematopoietic/Immune (0.240) Reproductive (0.120)	Cell Proliferative (0.640) Inflammation (0.440)	pINCY
80	595-639	Reproductive (0.211) Gastrointestinal (0.158) Urologic (0.158)	Cell Proliferative (0.684) Inflammation (0.263)	pINCY
81	425-469	Reproductive (0.222) Gastrointestinal (0.160) Nervous (0.148)	Cell Proliferative (0.568) Inflammation (0.259)	PSPORT1
82	774-818	Gastrointestinal (0.200) Hematopoietic/Immune (0.200) Nervous (0.200)	Cancer (0.600) Trauma (0.200) Inflammation (0.200)	PBLUESCRIPT
83	517-561	Nervous (0.526) Reproductive (0.132) Cardiovascular (0.105)	Cancer (0.342) Fetal (0.158) Inflammation (0.158)	PBLUESCRIPT
84	1944-1988	Nervous (0.250) Reproductive (0.188) Endocrine (0.125)	Cancer (0.438) Fetal (0.250) Trauma (0.250)	PSPORT1
85	1027-1071	Reproductive (0.219) Nervous (0.206) Hematopoietic/Immune (0.116)	Cancer (0.458) Inflammation (0.232) Fetal (0.181)	PSPORT1

1		3
	č	
	•	
,		`
-	3	2
:		Ć
t		٠

		TABLE 3 (cont.)		
Nucleotide Seq ID NO:	Selected Fragment	Tissue Expression (Fraction of Total)	Disease or Condition (Fraction of Total)	Vector
98	658-702	Reproductive (0.227) Hematopoietic/Immune (0.182) Gastrointestinal (0.167)	Cancer (0.424) Inflammation (0.318) Foral (0.167)	PT7T3
87	488-532	Nervous (0.200) Reproductive (0.200) Musculoskeletal (0.120)	Cancer (0.320) Fetal (0.320) Inflammation (0.320)	pincy
88	379-423	Cardiovascular (0.250) Nervous (0.250) Reproductive (0.250)	Cancer (0.417) Fetal (0.167) Neurological (0.167)	pINCY
68	632-676	Reproductive (0.417) Cardiovascular (0.167) Gastrointestinal (0.167)	Cancer (0.333) Fetal (0.167) Inflammation (0.167)	pINCY
06	258-302	Reproductive (0.294) Nervous (0.137) Hematopoietic/Immune (0.118)	Cancer (0.569) Fetal (0.431) Inflammation (0.176)	PINCY
91	433-477	Reproductive (0.750) Nervous (0.250)	Cancer (0.500) Inflammation (0.500)	PSPORT1
92	542-586	Gastrointestinal (0.273) Hematopoietic/Immune (0.273) Developmental (0.182)	Cancer (0.455) Inflammation (0.364) Fetal (0.182)	pINCY
63	797-817	Reproductive (0.272) Nervous (0.204) Cardiovascular (0.126)	Cancer (0.447) Inflammation (0.214) Fetal (0.155)	PSPORT1
94	541-585	Reproductive (0.273) Nervous (0.250) Cardiovascular (0.159)	Cancer (0.364) Fetal (0.205) Inflammation (0.205)	PSPORT1
95	111-155	Reproductive (0.250) Gastrointestinal (0.173) Nervous (0.154)	Cancer (0.481) Fetal (0.231) Inflammation (0.212)	PSPORT1
96	597-641	Reproductive (0.261) Cardiovascular (0.217) Nervous (0.130)	Cancer (0.391) Fetal (0.304) Inflammation (0.130)	PSPORT1

TABLE 3 (cont.)		
ABLE 3 (con	4	4
ABLE 3 (-	
ABLE	3	
ABL	•	
7		
7	-	į
Υ	2	
	<	ζ

Nucleotide				
Seg ID NO:	Selected Fragment	Tissue Expression (Fraction of Total)	Disease or Condition (Fraction of Total)	Vector
76	434-478	Cardiovascular (0.222) Endocrine (0.222)	Cancer (0.667) Fetal (0.111)	PSPORT1
		Gastrointestinal (0.222)	Neurological (0.111)	
	218-247	Reproductive (0.333)	Cancer (0.559)	
80	920-964	Gastrointestinal (0.129)	Inflammation (0.204)	PINCY
		Hematopoietic/Immune (0.118)	Fetal (0.183)	
	327-371	Gastrointestinal (0.211)	Cancer (0.421)	
66		Reproductive (0.211)	Fetal (0.316)	PINCY
		Cardiovascular (0.158)	Inflammation (0.158)	
	596-625	Reproductive (0.230)	Cancer (0.590)	
100		Nervous (0.164)	Inflammation (0.246)	PSPORT
		Gastrointestinal (0.131)	Fetal (0.082)	
	487-531	Cardiovascular (0.235)	Cancer (0.588)	
101		Reproductive (0.235)	Inflammation (0.176)	DINCY
		Hematopoietic/Immune (0.176)	Trauma (0.118)	
	218-247	Gastrointestinal (0.241)	Cancer (0.448)	
102	542-586	Hematopoietic/Immune (0.207)	Fetal (0.276)	DINCY
		Cardiovascular (0.138)	Inflammation (0.276)	
	219-263	Reproductive (0.500)	Cancer (0.500)	
103		Cardiovascular (0.250)	Inflammation (0.250)	DINCY
		Hematopoietic/Immune (0.250)	Trauma (0.250)	
	111-140	Hematopoietic/Immune (0.286)	Cancer (0.333)	
104	327-371	Nervous (0.238)	Fetal (0.286)	pINCY
		Reproductive (0.143)	Inflammation (0.286)	•
	243-281	Musculoskeletal (0.286)	Inflammation (0.429)	
105		Nervous (0.286)	Fetal (0.286)	DINCY
		Gastrointestinal (0.143)	Cancer (0.286)	
	271-315	Nervous (0.800)	Cancer (0.400)	
106		Reproductive (0.100)	Inflammation (0.200)	DINCY
		Cardiovascular (0.100)	Trauma (0.200)	

TARIE 3 (cont.)

		TABLE 3 (cont.)		
Nucleotide Seq ID NO:	Selected Fragment	Tissue Expression (Fraction of Total)	Disease or Condition (Fraction of Total)	Vector
107	489-533	Cardiovascular (0.364) Gastrointestinal (0.182) Reproductive (0.182)	Cancer (0.273) Trauma (0.273) Inflammation (0.182)	pINCY
108	156-200	Nervous (0.256) Reproductive (0.256) Hematopoietic/Immune (0.128)	Cancer (0.465) Fetal (0.291) Inflammation (0.186)	pincy
109	1459-1503	Cardiovascular (0.250) Hematopoietic/Immune (0.250) Nervous (0.167)	Inflammation (0.417) Cancer (0.333) Trauma (0.167)	PSPORT1
110	164-208	Nervous (1.000)	Neurological (1.000)	pINCY

FABLE 4

ــــــا	Nucleotide SEQ ID NO:	Library	Library Comment
	56	SPLNFET0 1	Library was constructed at Stratagene, using RNA isolated from a pool of fetal splean tissue. Pollowing vector packaging. 2x10° primary clones were then amplified to stabilize the library for long-term storage. Amplification may significantly skew sequence abundances.
	57	SYNORAB0 1	Library was constructed using RNA isolated from the synovial membrane tissue of a 68-year-old Caucasian female with rheumatoid arthritis.
	58	KIDNNOT0 1	Library was constructed using RNA isolated from the kidney tissue of a 64-year-old cureasian femmle, who died from an intracranial bleed. Patient history included 'rheumatoid arthritis and tobacco use.
	59	PLACNOB0 1	Library was constructed using RNA isolated from placenta.
	9	HNT2RAT0 1	Library was constructed at Stratagene (STR917211), using RNA isolated from the hNT2 cell line (derived from a human teratocarcinome that exhibited properties characteristic of a committed neuronal precursor). Cells were treated with retinoic acid for 24 hours.
75	61	EOSIHETO 2	Library was constructed using RNA isolated from peripheral blood cells apheresed from a 48-year-old Caucasian male. Patient history included hypercosinophilia. The cell population was determined to be greater than 77% eosinophils by Wright's stanning.
	62	HNT2AGT0 1	Library was constructed at Stratagene (STR937233), using RNA isolated from the hNT2 cell line (derived from a human teratocarcinoma that exhibited properties characteristic of a committed neuronal precursor). Cells were treated with retinoic acid for 5 weeks and with mitotic inhibitors for two weeks and allowed to mature for an additional 4 weeks in conditioned medium.
	63	PROSNOT0 2	Libbrary was constructed using RNA isolated from the diseased prostate tissue removed from a 50-year-old Coucasian male during a retropublic prostatectomy. Pathology indicated adenotibromatous hyperplasia was present. Pathology for the associated tumor tissue indicated adenocarcinoma Gleason grade 343. Patient history included dysurist, carcinoma in situ of prostate, coronary atherosclerosis, and hyperlipidemia.
	64	COLNTUTO 2	Library was constructed using RNA isolated from colon tumor tissue removed from a 75-year-old Gucasian male during a hemicolectomy. Pathology indicated invasive grade 3 adenocationma arising in a tubulovillous adenoma, which was distal to the lleocecal valve in the coun. The tumor penetrated deeply into the muscularis propria but not through it.

TABLE 4 (cont.)

Nuc1 SEO	Nucleotide SEO ID NO:	Library	Library Comment
	65	KIDNTUT0 1	Library was constructed using RNA isolated from the kidney tumor tissue removed from an 8-month-old female during nephrousterectomy. Pathology indicated Wilms' tumor (nephroblastome), which involved 30 percent of the renal parenchyma. Prior to surgery, the patient was receiving heparin anticoagulant therapy.
	99	THYRNOT0 3	Library was constructed using RNA isolated from thyroid tissue removed from the left thyroid of a 28-year-old Caucasian female during a complete thyroidectomy. Pathology indicated a small nodule of admonmatous hyperplasia present in the left thyroid. Pathology for the acconstant unor tissue indicated dominant follicular adenoma. Porning a well-encapsulated mass in the left thyroid.
	67	BLADTUTO 4	Library was constructed using RNA isolated from bladder tumor tissue removed from a 60-year-old Cueusaian male during a radical cystectomy, prostatectomy, and vasectomy. Pathology indicated grade 3 transitional cell carcinoma in the left bladder wall. Carcinoma in-situ was identified in the dome and trigone. Patient history included tobacco use. Family history included type I diabetes, a malignant neoplasm of the stomach, atherosclerotic coronary artery disease, and an acute myoraddial infarction.
1	89	PROSNOT1 5	Library was constructed using RNA isolated from diseased prostate tissue removed from a 66-year-old Caucasian male during radical prostatectomy and regional lymph node excision. Pathology indicated adenofibromatous hyperplasia. Pathology for the associated tumor tissue indicated an adenocarcinoma (Gleason grade 2+3). The parient presented with elevated prostate specific antigen (PSA). Family history included prostate cancer, and benigm hypertension.
	69	BRSTTUT0 8	Library was constructed using RNA isolated from breast tumor tissue removed from a 45-year-old Caucasian female during unilateral extended simple mastectomy. Pathology 167-year-old Caucasian female during unilateral extended simple mastectomy. Pathology form distanced invasive nuclear grade 2-3 adenocartinoma, ductal type, with 3 of 23 lymph nodes positive for metastatic disease. Oraster than 50% of the tumor volume was in situ, both comedo and non-comedo types. Immunostains were positive for estrogen/progesteron receptors, and uninvolved tissue showed prolifectative changes. The patient concurrently underwent a total abdominal hysterectomy. Patient history included valvuloplasty of mitral valve without replacement, rheumatic mitral insufficiency, and rheumatic heart diseases. Family history included acute myocardial infarction, atheorost-oronary artery disease, and type II diabetes.

PCT/US00/02237

BLE 4 (cont.)

į			
-	Nucleotide SEQ ID NO:	Library	Library Comment
	70	PROSNOT1 9	Library was constructed using RNA isolated from diseased prostate tissue removed from a 59-year-old caucasian male during a radical prostatectory with regional lymph mode excision. Pathology indicated adenotibromatous hyperplasia. Pathology for the associated tumor tissue indicated an adenocarcinoma (Glasson grade 3+3). The patient presented with elevated prostate-specific antigen (PSA). Patient history included colon diverticuli, asbestosis, and thrombophlabitis. Previous surgeries included a patial colectowy. Family history included benign hypertension, multiple myeloma, hyperligidemia and rheumatoid arthritis.
	71	THP1NOT0	Library was constructed using RNA isolated from untreated THP-1 cells. THP-1 (ATCC TIB 202) is a human promonocyte line derived from the peripheral blood of a 1-year-old Caucasian male with acute monocytic leukemia (ref: Int. J. Cancer (1980) 26:171).
7	72	THP1NOT0	Library was constructed using RNA isolated from untreated THP-1 cells. THP-1 (ATCC TIB 202) is a human promonocyte line derived from the peripheral blood of a 1-year-old Caucasian male with acute monocytic leukemia (ref: Int. J. Cancer (1980) 6:171).
	73	HIPOAZTO 1	Library was constructed from RNA isolated from diseased hippocampus tissue removed from the brain of a 74-year-old Caucasian male who died from Alzheimer's disease.
	74	HIPOAZTO 1	Library was constructed from RNA isolated from diseased hippocampus tissue removed from the brain of a 74-year-old Caucasian male who died from Alzheimer's disease.
	75	BRSTNOT1 3	Library was constructed using RNA isolated from breast tissue removed from the left medial lateral breast of a 36-year-old Goucaian famel dutring bilateral simple mastectomy and total breast reconstruction. Pathology indicated benign breast tissue. Patient history included a breast neoplasm, depressive disorder. Hyperlipidemia, chronic stomed unleer, and an ectopic pregnancy. Pamily history included myocardial inferction, cerebrovascular disease, atherosclerotic coronary artery disease, hyperlipidemia, skin cancer, breast cancer, depressive disorder, esophageal cancer, bone cancer, Hodgkin's lymphoma, bladder cancer, and heart condition.
	76	NPOLNOT0 1	Library was constructed using RNA isolated from nasal polyp tissue removed from a 78-year-old Gaucasian male during a nasal polypectowy. Pathology indicated a nasal polype coor, problem and nasal polype.

TABLE 4 (cont.)

Ĺ			
-	Nucleotide SEQ ID NO:	Library	Library Comment
	77	ADRENOT0 9	Library was constructed using RNA isolated from left adrenal gland tissue removed from a 43-year-old Caucasian male during nephrouseterectomy, regional lymph node excision, and unilateral left adrenalectomy. Pathology indicated no diagnostic abnormalities of the adrenal gland. Pathology for the associated tumor tissue indicated a grade 2 renal call carcinome mass in the posterior lower pole of the left kidney with invasion into the renal pairs.
	78	SPLNNOT1	Library was constructed using RNA isolated from diseased splean tissue removed from a 14-year-old Asian male during a total splenetromy. Pathology indicated changes consistent with idiopathic thrombocytopenic purpura. The patient presented with bruising.
78	79	CONCNOTO 1	Library was constructed using RNA isolated from chest wall soft tissue removed from a 63-year-old Caucasian male during a chest wall leasion destruction. Pathology indicated surgical margins were free of tumor. Pathology for the associated tumor tissue indicated invasive grade 3 adenocarcinoma, forming a mass that extended through the visceral pleura to involve parietal pleura. Patient history included MENA (multiple endocrine neoplasia) syndrome type I, abnormal secretion of gastrin, hemorrhage, that cook and allow the standard calculus of the kidney. Family history included prostate cancer, benign hypertension, stroke, atherosoclerotic coronary artery disease, type II diabetes, hyperlibidemia, and cancer of an unspecified location.
L	80	TONSNOT0	Library was constructed using RNA isolated from diseased left tonsil tissue removed from a 6-year-old Gaucasian male during adencionsillactomy. Pathology indicated reactive lymphoid hyperplasia, bilaterally. Pamily history included benign Mypertension, myocardial infarction, and atherosclerotic coronary artery disease
	81	LUNGNON0 3	This normalized library was constructed from 2.56 x 10° independent clones from a lung tissue library. NRA was made from lung tissue remaved from the left lobe a 59-year-old Gaucasian male during a segmental lung tissue remaved from the left lobe a 59-year-old Gaucasian male during a segmental lung resection. Pathology for the associated tumor tissue indicated a meatastatic grade 3 lof 4) osteosarcoma. Patient history included soft tissue cancer secondary cancer of the lung, prostate cancer and an acute duodenal ulcer with hemorrhage. Patient also received radiation therapy to the retroperiorinneum. Panily history included prostate cancer, breast cencer, and soute leavenia. The normalization and hybridization conditions were adapted from Sozes et al., PMSK (1994) 91:2223; Swarcop et al., NRR (1991) 19:1954; and Bonaldo et al., Genome Research (1996) 6:791

WO 00/44900

L	Marine San Allen		
	SEQ ID NO:	Library	Library Comment
	82	BMARNOT0 2	The library was constructed using Clontech RNA isolated from the bone marrow of 24 male and female Caucasian donors, 16 to 70 years old.
	83	SINTNOTO 2	The library was constructed using RNA isolated from the small intestine of a 55- year-old Caucasian female, who died from a subarachnoid hemorrhage. Serologies were positive for cytomegalouirus (CMV).
	84	CRBLNOT0 1	The library was constructed using RNA isolated from the cerebellum tissue of a 69- year-old Caucasian male who died from intronic obstructive pulmonary disease. Patient history included mycoralial infarction, hypertension, and osteoarthy-fre
	85	BRSTNOT0 3	The library was constructed using RNN isolated from diseased breest tissue removed from a \$4-year-old Caucasian femia deuting a bilateral radical mastectomy. Pathology for the associated tumor tissue indicated residual invasive grade mammary ductal adenocarcinoma. Patient history included kidney infection and condyloma ecuminatum. Pamily history included benign hypertension, hyperlipidemia and a malignant neoplasm of the colon.
79	98	BEPINON0 1	The normalized bronchial epithelium library was constructed from 5.12 million independent clones from a bronchial epithelium library. RNA was isolated from a bronchial epithelium primary cell line derived from a 64-year-old Caucasian male. The normalization and hybridization conditions were adapted from Soares et al., PNAS (1994) 91:9228, using a 24-hour reannealing hybridization period.
	87	THYRNOTO 3	The library was constructed using RNA isolated from thyroid tissue removed from the left thyroid of a 28-year-old teasasian female during a complete thyroididecromy. Pathology indicated a small module of adenomatous hyperplasia present in the left thyroid. Pathology for the associated turnor tissue indicated dominant follicular adenoma. Forming a well-encapsulated mass in the left thyroid.
	80	COLNFET0 2	The library was constructed using RNA isolated from the colon tissue of a Caucasian female fetus who died at 20 weeks' gestation.
	88	BLADNOT0 3	The library was constructed using RNA isolated from bladder tissue removed from an 80-year-old Caucasian female during a radical cyclectory and lymph node excision. Pathology for the associated tumor tissue indicated grade 3 invasive transitional cell carcinoma. Patient history included malignant neoplasm of the uterus, etheroscierosia, and atrial fibrillation. Family history included acute renal failure, osteoarchitis, and atherosclerosis.

PCT/US00/02237

TABLE 4 (cont.)

Nucleotide SEQ ID NO:	le Library	Library Comment
06	COLNNOT2	The library was constructed using RNA isolated from diseased colon tissue removed from a fe-year-old Caucasian male during a total colectomy with abdominal perineal resection. Pathology indicated gastritis and pencolonitis consistent with the acute phase of ulcerative colitis. Inflammation was more severe in the transverse colon with inflammation confined to the mucosa. The ascending and as signoid colon was mildly involved. Ramily history included irritable bowel syndrome.
91	BRSTNOT0	The library was constructed using RNA isolated from breast tissue removed from a 62- year-old East Indian female duting a unilateral extended simple mastectomy. Pathology for the associated tumor tissue indicated an invasive grade 3 ductal carcinoma. Patient history included benign hypertension, hyperlipidemia, and hematuria. Remily history included cerebrovascular and cardiovascular disease, hyperlipidemia, and livre cancer.
95	LIVRFET0	The library was constructed using RNA isolated from liver tissue removed from a Caucasian female fetus who died at 20 weeks' gestation. Family history included seven days of erythromycin treatment for bronchitis in the mother during the first trinsster.
93	BEPINOTO 1	The library was constructed using RNA isolated from a bronchial epithelium primary cell line derived from a 54-year-old Caucasian male.
94	BRAITUTO 2	The library was constructed using RNA isolated from brain tumor tissue removed from the frontal lobe of a 58-year-old Caucasian male during excision of a cerebral meningeal lesion Pathology indicated a grade 2 metastatic hyperrephrona. Patient history included a grade 2 renal call carcinoma, insomnia, and chronic airway obstruction. Pamily history included a malignant neoplasm of the kidney.
95	BRAITUTO 3	The library was constructed using RNA isolated from brain tumor tissue removed from the left frontal lobe a 17-year-old Caucasian female during excision of a cerebral meningeal lesion. Pathology indicated a grade 4 fibrillary giant and small-cell astrocytoma. Family history included benign hypertension and cerebrovascular disease.
96	BRSTTUTO 2	The library was constructed using RNA isolated from breast tumor tissue removed from a 54-year-old Gausaian femia duting a bilateral radical masterctony with reconstruction. Pathology indicated residual invasive grade 3 mammary ductal adenocarcinoma. The remaining breast parenchyma exhibited proliferative fibrocystic changes without artypia. One of 10 axillary lymph nodes had mestateic tumor. Patient history included kidney infection and condyloma acuminatum. Family history included henign hypertension, hyperlipidemia, and a malignant colon neoplasm.

TABLE 4 (cont.)

L			
	Nucleotide SEQ ID NO:	Library	Library Comment
	97	BRSTTUTO 2	The library was constructed using RNA isolated from breast tumor tissue removed from a S4-year-old Gacasian female during a bilateral radical mastectomy with reconstruction. Pathology indicated residual invasive grade 3 mammary ductal adenocarcinoma. The remaining breast parenchyma exhibited proliferative fibrocystic changes without atypia. One of 10 axillary lymph nodes had metastatic tumor. Bettent history included kidney infection and condyloma exuminatum. Femily history included benign hypertension, hyperalial and a metalial procytical decided and hypertension.
	86	BRSTNOT0	The library was constructed using RNN isolated from diseased breast tissue removed from a 43-year-old Caucasian female during a unilateral extended simple mastectomy. Pathology indicated mildly proliferative fibrocystic changes with epithelial hyperplasia, papillomatosis, and duct eiversia. Pathology for the associated tumor tissue indicated invasive grade 4, nuclear grade 3 mammary adenocarcinom with extensive comed necrosis. Family history included epilepsy, cardiovascular disease, and type II dispetes.
8	66	BRAINOT0 9	The library was constructed using RNA isolated from brain tissue removed from a Caucasian male fetus who died at 23 weeks, nestation
1	100	THP1AZS0 8	The library was constructed using RRM isolated from 5.76 million clones from a 5-aza-2"deoxyvgitdine treated THP-1 cell library. The library was subjected to libraricative hybridization using 5 million clones from an untreated THP-1 cell library. Hybridization conditions were adapted from Swarcop et al., NaR (1991) 19:1954; and Bonaldo et al., Genome Research (1996) 6:791. THP-1 (ATCC TIB 202) is a human promonocyte cell lihe derived from peripheral blood of a 1-year-old Caucasian male with acute monocytic lankamia
	101	LUNGTUT1 2	The library was constructed using RNA isolated from tumorous lung tissue removed from a 70-year-old Caucasian female during a lung lobectomy of the left upper lobe. Pathology indicated grade 3 (of 4) adenocarcinoma and vascular invasion. Patient history included tobacco abuse, depressive disorder, anxiety state, and skin cancer. Family history included cerebrovascular dieses, congestive heart failure, colon cancer, depressive disorder, and primary lives.
	102	HEAONOTO 5	The library was constructed using RNA isolated from aortic tissue removed from a 17-year-old Hispanic female who died from a quancher wound
	103	HEAONOTO 5	The library was constructed using RNA isolated from actic tissue removed from a 17- year-old Hispanic female who died from a gunshot wound.

PCT/US00/02237

ABLE 4 (cont.)

L	Nucleotide	Library	Library Comment
	104	EPIGNOT0 1	The library was constructed using RNA isolated from epiglottic tissue removed from a 71-year-old male during laryngectomy with right parathyroid biopsy. Pathology for the associated tumor tissue indicated recurrent grade 1 papillary thyroid carcinoma.
<u> </u>	105	CONCNOTO 1	The library was constructed using RNM isolated from chest wall soft tissue removed from a 61-year-old Caucasian male during a chest wall lession destruction. Pathology from the associated tumor tissue indicated invasive grade 3 adenocarcinoma forming a mass that extended through the visceral pleura to involve parietal pleura. Patient history included multiple endocrine neoplasia syndrom exper, chronic stomach upcartin, alcohol and tobacco abuse, calcium metabolism disease, chronic stomach ulcer with hemorrhage, lung cencer, and calculus of the kidney. Femily history included prostate cancer, benign hypertension, stroke, atheroscleroic coronary artery disease, type II diabetes. Apparlipidemia, and an unspecified cencer.
<u> </u>	106	BRONNOT0 1	The library was constructed using RNA isolated from bronchial tissue removed from a 15-year-old Caucasian male.
L	107	BRONNOT0 1	The library was constructed using RNA isolated from bronchial tissue removed from a 15-year-old Caucasian male.
2	108	LIVRNOT0 3	The library was constructed using RNA isolated from liver tissue removed from a Caucasian male fetus who died from Patau's syndrome (trisomy 13) at 20 weeks' pestation.
<u> </u>	109	LUNGNON0	The normalized library was constructed from 2.56 million independent clones from a lung fissue library. RRA was foolated from lung tissue removed from the left lobe as lung fissue library. RRA was foolated from lung tissue removed from the left lobe associated tumor tissue indicated a metastatic grade 3 (of 4) osteosarcoma. Patient history included soft tissue canner, secondary cancer of the lung, prostate cancer, accure duodenal ulcer with hemorrhaqe, and radiation therapy to the retroperitoneum. Family history included prostate cancer, breast cancer, and acute leukemia. The normalization and hybridization conditions were adapted from Soares et al., PMNS (1994) 91:9228; Swarcop et al., NAR (1991) 19:1954; and Bonaldo et al., Genome Research (1996) 6:991.
L	110	BRAIHCT0 2	The library was constructed using RNA isolated from diseased choroid plexus tissue removed from the brain of a 57-year-old Caucasian male who died from a cerebrovascular accident. Patient history included Huntington's disease and emphysema.

82

PCT/US00/02237

le 5

Program ABI FACTURA	Description A program that removes vector sequences and masks ambiguous bases in nucleic acid sequences.	Reference Perkin-Elmer Applied Biosystems, Foster City, CA.	Parameter Threshold
ABI/PARACEL FDF	A Fast Data Finder useful in comparing and annotating amino acid or nucleic acid sequences.	Perkin-Elmer Applied Biosystems, Foster City, CA; Paracel Inc., Pasadena, CA.	Mismatch <50%
ABI AutoAssembler	A program that assembles nucleic acid sequences.	Perkin-Elmer Applied Biosystems, Foster City, CA.	
BLAST	A Basic Local Alignment Search Tool use ful in sequence similarity search for amino acid and nucleic acid sequences. BLAST includes five functions: blastp, blastn, blasts, tibisstn, and tobasts.	Altschul, S.F. et al. (1990) J. Mol. Biol. 215:403-410; Altschul, S.F. et al. (1997) Nucleic Acids Res. 25: 3389-3402.	ESTs: Probability value= 1.0E-8 o less Full Length sequences: Probability value= 1.0E-10 or less
	A Pearson and Lipman algorithm that searches for similarity between a query sequence and a group of sequences of the same type. F ASTA comprises a least five functions: fasta, flasta, fastx, tlastx, and ssearch.	Pearson, W.R. and D.J. Lipman (1988) Proc. Natl Acade Sci. 8522446; Pearson, W.R. (1990) Mathods Enzymol. 183: 63-98, and Smith, T.F. and M. S. Waterman (1981) Adv. Appl. Math. 2-482-489.	Assembled ESTs: fasta E value=1.06E-6 Assembled ESTs: fasta Identity= 95% or greater and a state in the state
BLIMPS	A BLocks IMProved Searcher that matches a sequence against those in BLOCKS, PRINTS, DOMO, PRODOM, and PFAM databases to search for gene families, sequence homology, and structural fingerprint regions.	Henikoff, S and J.G. Henikoff, Nucl. Acid Res., 195365-72, 1991, J.G. Henikoff and S. Henikoff (1996) Methods Enzymol. 266.88- 105; and Attwood, T.K. et al. (1997) J. Chem. Inf. Comput. Soi. 37: 417-424.	Score=1000 or greater; Ratio of Score/Strength = 0.75 or larger; and, if applicable, Probability value= 1.0E-3 or less
IIMMER	An algorithm for searching a query sequence against hidden Markov model (HMM)-based databases of protein family consensus sequences, such as PFAM.	Krogh, A. et al. (1994) J. Mol. Biol., 235:1501-1531; Sonnhammer, E.L.L. et al. (1988) Nucleic Acids Res. 26:320-322.	Score = 10-50 bits for PFAM hits, depending on individual protein families

Table 5 (cont.)

Program	Description	Reference	Parameter Threshold
ProfiteScan	An algorithm that searches for structural and sequence motifs in protein sequences that match sequence patterns defined in Prosite.	Gribskov, M. et al. (1988) CABIOS 4:61-66; Gribskov, et al. (1989) Methods Enzymol. 183:146-159; Bairoch, A. et al. (1997) Nucleic Acids Res. 25: 217-221.	Normalized quality scores GCG- specified "HIGH" value for that particular Prosite motif. Generally, score=1,4-2.1.
Phred	A base-calling algorithm that examines automated sequencer traces with high sensitivity and probability.	Ewing, B. et al. (1998) Genome Res. 8:175-185; Ewing, B. and P. Green (1998) Genome Res. 8:186- 194.	
Рћар	A Phils Revised Assembly Program including SWAT and CrossMatch, programs based on efficient implementation of the Smith-Waterman algorithm, useful in searching sequence homology and assembling DNA sequences.	Smith, T.F. and M. S. Waterman (1981) Adv. Appl. Math. 2-482-489; Smith, T.F. and M. S. Waterman (1981) J. Mol. Biol. 147:195- 197; and Green, P., University of Washington, Seattle, WA.	Score= 120 or greater; Match length= 56 or greater
Consed	A graphical tool for viewing and editing Phrap assemblies	Gordon, D. et al. (1998) Genome Res. 8:195-202.	
SPScan	A weight matrix analysis program that scans protein sequences for the presence of secretory signal peptides.	Nielson, H. et al. (1997) Protein Engineering 10:1-6; Claverie, J.M. and S. Audic (1997) CABIOS 12: 431-439.	Score=3.5 or greater
Motifs	A program that searches amino acid sequences for patterns that matched those defined in Prosite.	Bairoch et al. <u>supra</u> ; Wisconsin Package Program Manual, version 9, page MS1-59, Genetics Computer Group, Madison, WI.	

WO 00/44900

PCT/US00/02237

What is claimed is:

20

- I. An isolated polypeptide comprising:
- a) an amino acid sequence selected from the group consisting of SEQ ID NO:1, SEQ ID
 NO:3-5, SEQ ID NO:7-14, SEQ ID NO:16-31, SEQ ID NO:33-34, SEQ ID NO:36-40, SEQ ID
 NO:42-48, SEQ ID NO:50-55.
 - b) a naturally occurring amino acid sequence having at least 90% sequence identity to an amino acid sequence selected from the group consisting of SEQ ID NO:1, SEQ ID NO:3-5, SEQ ID NO:7-14, SEQ ID NO:16-31, SEQ ID NO:33-34, SEQ ID NO:36-40,SEQ ID NO:42-48, SEQ ID NO:50-55,
 - c) a biologically active fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1, SEQ ID NO:3-5, SEQ ID NO:7-14, SEQ ID NO:16-31, SEQ ID NO:33-34, SEQ ID NO:36-40, SEQ ID NO:42-48, SEQ ID NO:50-55, or
- d) an immunogenic fragment of an amino acid sequence selected from the group consisting of SEQ ID NO:1, SEQ ID NO:3-5, SEQ ID NO:7-14, SEQ ID NO:16-31, SEQ ID NO:33-34, SEQ ID NO:36-40,SEQ ID NO:42-48, SEQ ID NO:50-55.
 - An isolated polypeptide of claim 1, having an amino acid sequence selected from the group consisting of SEQ ID NO:1, SEQ ID NO:3-5, SEQ ID NO:7-14, SEQ ID NO:16-31, SEQ ID NO:33-34, SEQ ID NO:36-40,SEQ ID NO:42-48, SEQ ID NO:50-55.
 - 3. An isolated polynucleotide encoding a polypeptide of claim 1.
- An isolated polynucleotide of claim 3, having a sequence selected from the group
 consisting of SEQ ID NO:56-110.
 - A recombinant polynucleotide comprising a promoter sequence operably linked to a polynucleotide of claim 3.
- 6. A cell transformed with a recombinant polynucleotide of claim 5.
 - 7. A transgenic organism comprising a polynucleotide of claim 5.
 - 8. A method for producing a polypeptide of claim 1, the method comprising:
- a) culturing a cell under conditions suitable for expression of the polypeptide, wherein said

WO 00/44900 PCT/US00/02237

cell is transformed with a recombinant polynucleotide, and said recombinant polynucleotide comprises a promoter sequence operably linked to a polynucleotide encoding the polypeptide of claim 1, and

b) recovering the polypeptide so expressed.

5

10

15

35

a come as a lake was a state of a sure and a stage

- 9. An isolated antibody which specifically binds to a polypentide of claim 1.
- 10. An isolated polynucleotide comprising:
- a) a polynucleotide sequence selected from the group consisting of SEQ ID NO:56-110,
- a naturally occurring polynucleotide sequence having at least 90% sequence identity to a polynucleotide sequence selected from the group consisting of SEQ ID NO:56-110,
 - c) a polynucleotide sequence complementary to a), or
 - d) a polynucleotide sequence complementary to b).
- An isolated polynucleotide comprising at least 60 contiguous nucleotides of a polynucleotide of claim 10.
- A method for detecting a target polynucleotide in a sample, said target polynucleotide
 having a sequence of a polynucleotide of claim 10, the method comprising:
 - a) hybridizing the sample with a probe comprising at least 16 contiguous nucleotides comprising a sequence complementary to said target polynucleotide in the sample, and which probe specifically hybridizes to said target polynucleotide, under conditions whereby a hybridization complex is formed between said probe and said target polynucleotide, and
- 25 b) detecting the presence or absence of said hybridization complex, and, optionally, if present, the amount thereof.
 - 13. A method of claim 12, wherein the probe comprises at least 30 contiguous nucleotides.
- 30 14. A method of claim 12, wherein the probe comprises at least 60 contiguous nucleotides.
 - 15. A pharmaceutical composition comprising an effective amount of a polypeptide of claim 1 and a pharmaceutically acceptable excipient.
 - 16. A method of treating a disease or condition associated with decreased expression of

WO 00/44900

PCT/US00/02237

functional NuABP, comprising administering to a patient in need of such treatment the pharmaceutical composition of claim 15.

- 17. A method for screening a compound for effectiveness as an agonist of a polypeptide of claim 1, the method comprising:
 - a) exposing a sample comprising a polypeptide of claim 1 to a compound, and
 - b) detecting agonist activity in the sample.
- 18. A pharmaceutical composition comprising an agonist compound identified by a method 10 of claim 17 and a pharmaceutically acceptable excipient.
 - 19. A method of treating a disease or condition associated with decreased expression of functional NuABP, comprising administering to a patient in need of such treatment a pharmaceutical composition of claim 18.

15

- 20. A method for screening a compound for effectiveness as an antagonist of a polypeptide of claim 1, the method comprising:
 - a) exposing a sample comprising a polypeptide of claim 1 to a compound, and
 - b) detecting antagonist activity in the sample.

20

30

- 21. A pharmaceutical composition comprising an antagonist compound identified by a method of claim 20 and a pharmaceutically acceptable excipient.
- A method for treating a disease or condition associated with overexpression of functional
 NuABP, comprising administering to a patient in need of such treatment a pharmaceutical composition of claim 21.
 - 23. A method for screening a compound for effectiveness in altering expression of a target polynucleotide, wherein said target polynucleotide comprises a sequence of claim 4, the method comprising:
 - a) exposing a sample comprising the target polynucleotide to a compound, and
 - b) detecting altered expression of the target polynucleotide.

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 3 August 2000 (03.08.2000)

PCT

(10) International Publication Number WO 00/44900 A3

- C12Q 1/68, C07K 14/47, 16/18, G01N 33/68, A61K 38/17
- (21) International Application Number: PCT/US00/02237
- (22) International Filing Date: 28 January 2000 (28.01.2000)
- (25) Filing Language: English
- (26) Publication Language:

(51) International Patent Classification?:

English

C12N 15/12.

- (30) Priority Data: 60/117.905 29 January 1999 (29.01.1999) 60/117,904 29 January 1999 (29.01.1999)
- (63) Related by continuation (CON) or continuation-in-part (CIP) to earlier applications:

211 60/117.904 (CIP) 29 January 1999 (29,01,1999) Filed on HS 60/117,905 (CIP) Filed on 29 January 1999 (29.01.1999)

- (71) Applicant (for all designated States except US): INCYTE PHARMACEUTICALS, INC. [US/US]; 3174 Porter Drive, Palo Alto, CA 94304 (US).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): TANG, Y., Tom [CN/US]; 4230 Ranwick Court, San Jose, CA 95118 (US). LAL, Pretti [IN/US]; 2382 Lass Drive, Santa Clara, CA 95054 (US). HILLMAN, Jennifer, L. [US/US]; 230 Monrow Drive #12, Mountain View, CA 94040 (US). YUE, Henry [US/US]; 826 Lois Avenue, Sunnyvale, CA 94087 (US), AZIMZAI, Yalda [US/US]; 2045 Rock Springs Drive, Hayward, CA 94545 (US). LU, Aina, M., D. [US/US]; 55 Park Belmont Place, San Jose, CA 95136 (US). BAUGHN, Mariah, R. [US/US]; 14244

Santiago Road, San Leandro, CA 94577 (US), TRAN, Bao [US/US]; 744 Kiely Boulevard, Santa Clara, CA 95051 (US), SHIH, Leo, L. [US/US]; Apartment B., 1081 Tanland Drive, Palo Alto, CA 94303 (US), AU-YOUNG, Janice, L. [US/US]; 233 Golden Eagle Lane, Brisbane, CA 94005 (US).

- (74) Agents: HAMLET-COX, Diana et al.: Incyte Pharmaceuticals, Inc., 3174 Porter Drive, Palo Alto, CA 94304 (US).
- (81) Designated States (national): AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT. BE, CH. CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published:

- With international search report.
 - Before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments.
- (88) Date of publication of the international search report: 30 November 2000

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette

(54) Title: NUCLEIC-ACID BINDING PROTEINS

(57) Abstract: The invention provides human nucleic-acid binding proteins (NuABP) and polynucleotides which identify and encode NuABP. The invention also provides expression vectors, host cells, antibodies, agonists, and antagonists. The invention also provides methods for diagnosing, treating, or preventing disorders associated with expression of NuABP.

HERBERT BERTHAM BERTHAM

DECLARATION AND POWER OF ATTORNEY FOR UNITED STATES PATENT APPLICATION

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name, and

I believe that I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if more than one name is listed below) of the subject matter which is claimed and for which a United States patent is sought on the invention entitled

NUCLEIC-ACID BINDING PROTEINS

ne specification of which:	
/ is attached hereto.	
/_/ was filed on as application Serial No contains an X //, was amended on	_ and if this box
/X_/ was filed as Patent Cooperation Treaty international application No. anuary 28, 2000, this box contains an X /_/, was amended on under Patent Conticle 19 on 2001, and if this box contains an X /_/, was amended or	ooperation Treaty

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge my duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a).

I hereby claim the benefit under Title 35, United States Code, §119 or §365(a)-(b) of any foreign application(s) for patent or inventor's certificate indicated below and of any Patent Cooperation Treaty international applications(s) designating at least one country other than the United States indicated below and have also identified below any foreign application(s) for patent or inventor's certificate and Patent Cooperation Treaty international application(s) designating at least one country other than the United States for the same subject matter and having a filing date before that of the application for said subject matter the priority of which is claimed:

Country	Number	Filing Date	Priority Claimed
			/_/ Yes /_/ No // Yes // No
			/_/ Tes /_/ No

I hereby claim the benefit under Title 35, United States Code, §119(e) of any United States provisional application(s) listed below.

Application		Status (Pending,
Serial No.	Filed	Abandoned, Patented)
60/117,905	January 29, 1999	Expired
60/117,904	January 29, 1999	Expired

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in said prior application(s) in the manner required by the first paragraph of Title 35, United States Code §112, I acknowledge my duty to disclose material information as defined in Title 37 Code of Federal Regulations, §1.56(a) which occurred between the filing date(s) of the prior application(s) and the national or Patent Cooperation Treaty international filing date of this application:

Application		Status (Pending,
Serial No.	Filed	Abandoned, Patented)

I hereby appoint the following:

Lucy J. Billings	Reg. No. 36,749
Michael C. Cerrone	Reg. No. 39,132
Diana Hamlet-Cox	Reg. No. 33,302
Richard C. Ekstrom	Reg. No. 37,027
Barrie D. Greene	Reg. No. 46,740
Lynn E. Murry	Reg. No. 42,918
Shirley A. Recipon	Reg. No. 47,016
Susan K. Sather	Reg. No. 44,316
Michelle M. Stempien	Reg. No. 41,327
David G. Streeter	Reg. No. 43,168
Stephen Todd	Reg. No. 47,139
P. Ben Wang	Reg. No. 41,420

respectively and individually, as my patent attorneys and/or agents, with full power of substitution and revocation, to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith. Please address all communications to:

LEGAL DEPARTMENT NCYTE GENOMICS, INC. 3160 PORTER DRIVE, PALO ALTO, CA 94304

TEL: 650-855-0555

FAX: 650-849-8886 or 650-845-4166

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

	1-00	
First Joint Inventor:	Full name:	Y. Tom Tang
	Signature:	U. halos
	Date:	Sept. 10,2001
	Citizenship	United States of America
	Residence:	San Jose, California CA
	P.O. Address:	4230 Ranwick Court
		San Jose, California 95118
	2-00	
Second Joint Inventor:	Full name:	Preeti Lal
	Signature:	Prechi W
	Date:	September 10, 2001
	Citizenship	India
	Residence:	Santa Clara, California 🔾 😝
	P.O. Address:	P.O. Box 5142 Santa Clara, California 95056
		Danta Ciara, Cantorna 75050

5518 Boulder Canyon Dr. Castro Valley, California 94552

Docket No.: PF-0662 USN

	3-00	
Third Joint Inventor:	Full name:	Jennifer L. Hillman
	Signature:	hit Hall
	Date:	Sutley 21, 2001
	Citizenship	United States of America
	Residence:	Mountain View, California 🖙
	P.O. Address:	230 Monroe Drive, #17 Mountain View, California 94040
	4-00	
Fourth Joint Inventor:	Full name:	Henry Yue
	Signature:	Hanny hee
	Date:	September 34 , 2001
	Citizenship	United States of America
	Residence:	Sunnyvale, California CA
	P.O. Address:	826 Lois Avenue
		Sunnyvale, California 94087
	5-W	
Fifth Joint Inventor:	Full name:	Yalda Azimzai
	Signature:	Alda (mm zn)
	Date:	September 13, 2001
	Citizenship	United States of America
	Residence:	Castro Valley, California

P.O. Address:

	6-00	
Sixth Joint Inventor:	Full name:	Dyung Aina M. Lu
	Signature:	Sina lu
	Date:	Sept 7 , 2001
	Citizenship	United States of America
	Residence:	San Jose, California
	P.O. Address:	233 Coy Drive San Jose, California 95123
	7-00	
Seventh Joint Inventor:	Full name:	Mariah R. Baughn
	Signature:	Muck. Blu
	Date:	September 5, 2001
	Citizenship	United States of America
	Residence:	San Leandro, California 🗲
	P.O. Address:	14244 Santiago Road
		San Leandro, California 94577
Eighth Joint Inventor:	8 ∕∕∕ Full name:	Bao Tran
Eighth John Inventor.	Signature:	2007
	5	, 2001
	Date:	
	Citizenship	United States of America
	Residence:	Santa Clara, California
	P.O. Address:	750 Salberg Avenue

D9889616.D5D8DR

Brisbane, California 94005

Docket No.: PF-0662 USN

9-50 Ninth Joint Inventor: Full name: Leo L. Shih Signature: , 2001 Date: United States of America Citizenship East Palo Alto, California Residence: P.O. Address: 908 O'Connor St. East Palo Alto, California 94303 10-00 **Tenth Joint Inventor:** Full name: Janice Au-Young Signature: , 2001 Date: United States of America Citizenship Brisbane, California Residence: P.O. Address: 233 Golden Eagle Lane

DECLARATION AND POWER OF ATTORNEY FOR INITED STATES PATENT APPLICATION

As a below named inventor, I hereby declare that:

the specification of which:

My residence, post office address and citizenship are as stated below next to my name, and

I believe that I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if more than one name is listed below) of the subject matter which is claimed and for which a United States patent is sought on the invention entitled

NUCLEIC-ACID BINDING PROTEINS

// is attached hereto.
// was filed on as application Serial No and if this box contains an X //, was amended on
/X / was filed as Patent Cooperation Treaty international application No. PCT/US00/02237 of January 28, 2000, this box contains an X / /, was amended on under Patent Cooperation Treaty Article 19 on 2001, and if this box contains an X / /, was amended on
I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge my duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, \$1.56(a).

I hereby claim the benefit under Title 35, United States Code, §119 or §365(a)-(b) of any foreign application(s) for patent or inventor's certificate indicated below and of any Patent Cooperation Treaty international applications(s) designating at least one country other than the United States indicated below and have also identified below any foreign application(s) for patent or inventor's certificate and Patent Cooperation Treaty international application(s) designating at least one country other than the United States for the same subject matter and having a filing date before that of the application for said subject matter the priority of which is claimed:

=0

Country	Number	Filing Date	Priority Claimed
			/_/ Yes /_/ No
			// Yes // No

I hereby claim the benefit under Title 35, United States Code, §119(e) of any United States provisional application(s) listed below.

Application		Status (Pending,
Serial No.	Filed	Abandoned, Patented)
60/117,905	January 29, 1999	Expired
60/117,904	January 29, 1999	Expired

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in said prior application(s) in the manner required by the first paragraph of Title 35, United States Code §112, I acknowledge my duty to disclose material information as defined in Title 37 Code of Federal Regulations, §1.56(a) which occurred between the filing date(s) of the prior application(s) and the national or Patent Cooperation Treaty international filing date of this application:

Application		Status (Pending,	
Serial No.	Filed	Abandoned, Patented)	

I hereby appoint the following:

Lucy J. Billings	Reg. No. 36,749
Michael C. Cerrone	Reg. No. 39,132
Diana Hamlet-Cox	Reg. No. 33,302
Richard C. Ekstrom	Reg. No. 37,027
Barrie D. Greene	Reg. No. 46,740
Lynn E. Murry	Reg. No. 42,918
Shirley A. Recipon	Reg. No. 47,016
Susan K. Sather	Reg. No. 44,316
Michelle M. Stempien	Reg. No. 41,327
David G. Streeter	Reg. No. 43,168

respectively and individually, as my patent attorneys and/or agents, with full power of substitution and revocation, to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith. Please address all communications to:

LEGAL DEPARTMENT INCYTE GENOMICS, INC. 3160 PORTER DRIVE, PALO ALTO, CA 94304

TEL: 650-855-0555 F/

FAX: 650-849-8886 or 650-845-4166

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

First Joint Inventor:	Full name:	Y. Tom Tang	
	Signature:		
	Date:	, 2001	
	Citizenship	United States of America	
	Residence:	San Jose, California	
	P.O. Address:	4230 Ranwick Court San Jose, California 95118	
Second Joint Inventor:	Full name:	Preeti Lal	
	Signature:		
	Date:	, 2001	
	Citizenship	India	
	Residence:	Santa Clara, California	
	P.O. Address:	P.O. Box 5142	

5518 Boulder Canyon Dr. Castro Valley, California 94552

Docket No.: PF-0662 USN

Third Joint Inventor:	Full name:	Jennifer L. Hillman
	Signature: Date:	, 2001
	Citizenship	United States of America
	Residence:	Mountain View, California
	P.O. Address:	230 Monroe Drive, #17 Mountain View, California 94040
Fourth Joint Inventor:	Full name:	Henry Yue
Touris Joint Inventor.	Signature:	ittiny rue
	Date:	, 2001
	Citizenship	United States of America
	Residence:	Sunnyvale, California
	P.O. Address:	826 Lois Avenue Sunnyvale, California 94087
Fifth Joint Inventor:	Full name:	Yalda Azimzai
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
	Residence:	Castro Valley, California

P.O. Address:

Santa Clara, California95051

Docket No.: PF-0662 USN

C' d I ' I I		
Sixth Joint Inventor:	Full name:	Dyung Aina M. Lu
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
	Residence:	San Jose, California
	P.O. Address:	233 Coy Drive
		San Jose, California 95123
Seventh Joint Inventor:	Full name:	Mariah R. Baughn
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
	Residence:	San Leandro, California
	P.O. Address:	14244 Santiago Road
		San Leandro, California 94577
		1
Eighth Joint Inventor:	Full name:	Bao Tran
	Signature: 🚅	
	Date:	Set 26, 2001
	Citizenship	United States of America
	Residence:	Santa Clara, California
	P.O. Address:	750 Salberg Avenue

Ninth Joint Inventor:	Full name:	Leo L. Shih
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
	Residence:	East Palo Alto, California
	P.O. Address:	908 O'Connor St. East Palo Alto, California 94303
Tenth Joint Inventor:	Full name:	Janice Au-Young
	Signature:	-
	Date:	, 2001
	Citizenship	United States of America
	Residence:	Brisbane, California
	P.O. Address:	233 Golden Eagle Lane Brisbane, California 94005

DECLARATION AND POWER OF ATTORNEY FOR UNITED STATES PATENT APPLICATION

As a below named inventor, I hereby declare that:

the specification of which:

/ / is attached hereto.

My residence, post office address and citizenship are as stated below next to my name, and

I believe that I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if more than one name is listed below) of the subject matter which is claimed and for which a United States patent is sought on the invention entitled

NUCLEIC-ACID BINDING PROTEINS

	as application Serial No	
contains an X. //. was an	nended on	 '
January 28, 2000, this box	nt Cooperation Treaty international applic contains an X /_/, was amended on under	Patent Cooperation Treaty
Article 19 on 2001	, and if this box contains an X /_/, was an	mended on
•	have reviewed and understand the conter claims, as amended by any amendment re	
	duty to disclose information which is mate with Title 37, Code of Federal Regulations	

I hereby claim the benefit under Title 35, United States Code, §119 or §365(a)-(b) of any foreign application(s) for patent or inventor's certificate indicated below and of any Patent Cooperation Treaty international applications(s) designating at least one country other than the United States indicated below and have also identified below any foreign application(s) for patent or inventor's certificate and Patent Cooperation Treaty international application(s) designating at least one country other than the United States for the same subject matter and having a filing date before that of the application for said subject matter the priority of which is claimed:

Country	Number	Filing Date	Priority Claimed
			/_/ Yes /_/ No
			/_/ Yes /_/ No

I hereby claim the benefit under Title 35, United States Code, §119(e) of any United States provisional application(s) listed below.

Application		Status (Pending,
Serial No.	Filed	Abandoned, Patented)
60/117,905	January 29, 1999	Expired
60/117,904	January 29, 1999	Expired

I hereby claim the benefit under Title 35. United States Code, §120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in said prior application(s) in the manner required by the first paragraph of Title 35, United States Code §112, I acknowledge my duty to disclose material information as defined in Title 37 Code of Federal Regulations, §1.56(a) which occurred between the filing date(s) of the prior application(s) and the national or Patent Cooperation Treaty international filing date of this application:

Application		Status (Pending,
Serial No.	Filed	Abandoned, Patented)

I hereby appoint the following:

Lucy J. Billings	Reg. No. 36,749
Michael C. Cerrone	Reg. No. 39,132
Diana Hamlet-Cox	Reg. No. 33,302
Richard C. Ekstrom	Reg. No. 37,027
Barrie D. Greene	Reg. No. 46,740
Lynn E. Murry	Reg. No. 42,918
Shirley A. Recipon	Reg. No. 47,016
Susan K. Sather	Reg. No. 44,316
Michelle M. Stempien	Reg. No. 41,327
David G. Streeter	Reg. No. 43,168
P. Ben Wang	Reg. No. 41,420

respectively and individually, as my patent attorneys and/or agents, with full power of substitution and revocation, to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith. Please address all communications to:



LEGAL DEPARTMENT INCYTE GENOMICS, INC. 3160 PORTER DRIVE, PALO ALTO, CA 94304

TEL: 650-855-0555 FAX: 650-849-8886 or 650-845-4166

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

First Joint Inventor:	Full name:	Y. Tom Tang
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
	Residence:	San Jose, California
	P.O. Address:	4230 Ranwick Court San Jose, California 95118
Second Joint Inventor:	Full name:	Preeti Lal
	Signature:	
	Date:	, 2001
	Citizenship	India
	Residence:	Santa Clara, California
	P.O. Address:	P.O. Box 5142 Santa Clara, California 95056

Valley, California 94552

Docket No.: PF-0662 USN

Third Joint Inventor:	Full name:	Jennifer L. Hillman
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
	Residence:	Mountain View, California
	P.O. Address:	230 Monroe Drive, #17 Mountain View, California 94040
Fourth Joint Inventor:	Full name:	Henry Yue
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
<i>:</i>	Residence:	Sunnyvale, California
	P.O. Address:	826 Lois Avenue Sunnyvale, California 94087
· ·		
Fifth Joint Inventor:	Full name:	Yalda Azimzai
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
	Residence:	Castro Valley, California
	P.O. Address:	5518 Boulder Canyon Dr. Castro

Sixth Joint Inventor:	Full name: Signature: Date: Citizenship Residence: P.O. Address:	, 2001 United States of America San Jose, California 233 Coy Drive
		San Jose, California 95123
Seventh Joint Inventor:	Full name: Signature:	Mariah R. Baughn
	Date:	, 2001
	Citizenship	United States of America
	Residence:	San Leandro, California
	P.O. Address:	14244 Santiago Road San Leandro, California 94577
Eighth Joint Inventor:	Full name:	Bao Tran
	Signature:	
	Date:	, 2001
	Citizenship	United States of America
	Residence:	Santa Clara, California
	P.O. Address:	750 Salberg Avenue Santa Clara, California95051

Ninth Joint Inventor:

Full name: Signature: Leo L. Shih

, 2001

, 2001

Date:

october 27 United States of America

Citizenship

East Palo Alto, California

Residence: P.O. Address:

908 O'Connor St.

East Palo Alto, California 94303

Tenth Joint Inventor:

Full name:

Janice Au-Young

Signature:

Date: Citizenship

United States of America

Residence:

Brisbane, California

P.O. Address:

233 Golden Eagle Lane

Brisbane, California 94005

SEQUENCE LISTING

```
<110> INCYTE PHARMACEUTICALS, INC.
       TANG, Y. Tom
       LAL, Preeti
       HILLMAN, Jennifer L.
       YUE, Henry
       AZIMZAI, Yalda
       LU, Aina M.D.
       BAUGHN, Mariah R.
       TRAN, Bao
       SHIH, Leo L.
       AU-YOUNG, Janice
<120> NUCLEIC ACID-BINDING PROTEINS
<130> PF-0662 PCT
<140> To Be Assigned
<141> Herewith
<150> 60/117,905; 60/117,904
<151> 1999-01-29: 1999-01-29
<160> 110
<170> PERL Program
<210> 1
<211> 754
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 025733CD1
<400> 1
Met Ala Ala Gly Ser Arg Lys Arg Arg Leu Ala Glu Leu Thr
  1
                                      10
Val Asp Glu Phe Leu Ala Ser Gly Phe Asp Ser Glu Ser Glu Ser
                  20
                                      25
                                                           30
Glu Ser Glu Asn Ser Pro Gln Ala Glu Thr Arg Glu Ala Arg Glu
                 35
                                      4 O
                                                           45
Ala Ala Arg Ser Pro Asp Lys Pro Gly Gly Ser Pro Ser Ala Ser
                                      55
Arg Arg Lys Gly Arg Ala Ser Glu His Lys Asp Gln Leu Ser Arg
                  65
                                      70
Leu Lys Asp Arg Asp Pro Glu Phe Tyr Lys Phe Leu Gln Glu Asn
                 80
                                      85
                                                          90
Asp Gln Ser Leu Leu Asn Phe Ser Asp Ser Asp Ser Ser Glu Glu
                 95
                                     100
Glu Glu Gly Pro Phe His Ser Leu Pro Asp Val Leu Glu Glu Ala
                 110
                                     115
                                                         120
Ser Glu Glu Glu Asp Gly Ala Glu Glu Gly Glu Asp Gly Asp Arg
                 125
                                     130
Val Pro Arg Gly Leu Lys Gly Lys Lys Asn Ser Val Pro Val Thr
                140
                                     145
                                                         150
Val Ala Met Val Glu Arg Trp Lys Gln Ala Ala Lys Gln Arg Leu
```

155

1. 1. 21 1 97 1.

170

PCT/US00/02237

Val Ala Thr Thr Arg Glv Asp Gln Glu Ser Ala Glu Ala Asn Lvs 185 190 195 Phe Phe Gln Val Thr Asp Ser Ala Ala Phe Asn Ala Leu Val Thr 210 200 205 Gly Cys Ile Arg Asp Leu Ile Gly Cys Leu Gln 215 220 225 Ser Ala Lvs Asp Ser Ser Arg Met Leu Gln Pro Ser Ser 230 235 240 Pro Leu Trp Gly Lys Leu Arg Val Asp Ile Lys Ala Tyr Leu Gly 245 255 250 Leu Ser Ala Ile Gln Leu Val Ser Cvs Leu Ser Glu Thr Thr Val 260 265 His Ile Ser Val Leu Val Pro Cvs Phe Leu Ala Val Leu Arg 275 280 285 Pro Lys Gln Val Cys Arg Met Leu Leu 290 295 300 Tro Ser Thr Glv Leu Glu Glu Ser Leu Arg 315 305 310 Cys Arg His Lys Lys Glv Leu Ser Arg Va1 Asp 330 320 325 Leu Lys Gln Met Tyr Ile Thr Tyr Val Arg Asn Cys Lvs 335 340 345 Trp Ser Pro Gly Ala Leu Pro Phe Tle Ser Phe Met Gln 350 355 Thr Glu Leu Leu Ala Leu Glu Pro Gln 365 370 His Ala Phe Leu Tyr Ile Arg Gln Leu Ala Ile His Leu Arg 380 385 390 Lys Lys Glu Thr Tyr Gln Ser Val Tyr Thr Thr Arg Asn 395 400 405 Leu Trp Cys Arg Val Leu Ser Tvr Val His Cys Leu Phe 415 410 Thr Ala Glv Pro Ser Glu Ala Leu Gln Pro Leu 425 430 Gln Val Ile Ile Gly Cys Ile Lys Leu 440 445 450 Tyr Pro Leu Arg Met His Cys Ile Arg Ala Leu Thr Leu Leu 455 460 Ser Gly Ser Ser Gly Ala Phe Ile Pro Val Len 470 475 Met Glu Met Phe Gln Gln Val Asp Phe Asn Arg Lys 495 485 490 Ser Ser Lys Pro Ile Asn Phe Ser Val Ile Leu Lys Leu Ser Asn 500 505 510 Gln Asn Leu Gln Glu Lys Ala Tyr Arg Asp Gly Leu Val Glu 520 515 Leu Tyr Asp Leu Thr Leu Glu Tyr Leu His Cys 540 530 535 Ile Gly Phe Pro Glu Leu Val Leu Pro Val Val Leu Gln Leu Lys 545 550 555 Phe Leu Arg Glu Cys Lys Val Ala Asn Tyr Cys Arg Gln Val 560 565 570 Gln Gln Leu Leu Gly Lys Val Gln Glu Asn Cys 575 580 585 Val Ser Phe Gly Val Ser Glu Gln Gln Ala Ser Arg Arg Gln Arg 590 595 600

Val Glu Ala Trp Glu Lys Leu Thr Arg Glu Glu Gly Thr Pro Leu

605

Thr Pro Lys Leu Phe His Glu Val Val Gln Ala Phe Arg Ala Ala

175

PCT/US00/02237

Thr Leu Tyr Tyr Ser His Trp Arg Lys Leu Arg Asp Arg Glu Ile 620 625 630 Ser Gly Lys Glu Arg Leu Glu Asp Leu Asn Phe Gln Leu Glu Tle 635 640 Pro Glu Ile Lys Arg Arg Lys Met Ala Asp 650 655 660 Arg Lys Gln Phe Lys Asp Leu Phe Asp Leu Asn Ser Ser Glu Glu 665 670 675 Asp Asp Thr Glu Gly Phe Ser Glu Arg Gly Ile Leu Arg Pro Leu 680 685 Ser Thr Arg His Glv Val Glu Asp Asp Glu Glu Asp Glu Glu Glu 700 705 695 Gly Glu Glu Asp Ser Ser Asn Ser Glu Gly Glu Trp Ser Trp Asp 710 715 720 Ala Glu Ala Gly Leu Ala Pro Gly Glu Leu Gln Gly Asp Pro Asp 725 730 735 Gln Leu Ala Gln Gly Pro Glu Asp Glu Leu Glu Asp Leu Gln Leu 740 745 750 Ser Glu Asp Asp

```
<210> 2
<211> 593
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 079702CD1
```

<400> 2 Met Arg Asp Ser Thr Gly Ala Gly Asn Ser Leu Val His Lys Arg 1 5 15 Arg Asn Gln Lys Thr Pro Thr Ser Leu Thr Lys Ser Pro Leu Ara 20 25 Leu Ser Leu Gln Asp Gly His Lys Ala Lys 40 35 45 Phe Glu Glu Gly Gln Asp Val Leu Ala Arg Trp Ser Asp Gly Leu 55 60 Ile Lys Lys Ile Asn Ile Leu Lys Gln Ser Phe Tyr Leu Gly Thr 65 70 75 Cys Phe Ile Ile Phe Glu Asp Ser Ser Lys Ser Trp Val Leu Trp 80 85 90 Lys Asp Ile Gln Thr Gly Ala Thr Gly Ser Gly Glu Met Cys 95 100 105 Thr Ile Cys Gln Glu Glu Tyr Ser Glu Ala Pro Asn Glu Met Val 110 115 120 Ile Cys Asp Lys Cys Gly Gln Gly Tyr His Gln Leu Cys His Thr 125 130 135 Pro His Ile Asp Ser Ser Val Ile Asp Ser Asp Glu Lys Trp Leu 140 145 150 Cys Arg Gln Cys Val Phe Ala Thr Thr Thr Lys Arg Gly Gly Ala 160 165 Leu Lys Lys Gly Pro Asn Ala Lys Ala Leu Gln Val Met Lys Gln 170 175 180 Thr Leu Pro Tyr Ser Val Ala Asp Leu Glu Trp Asp Ala Gly His 195 185 190 Lys Thr Asn Val Gln Gln Cys Tyr Cys Tyr Cys Gly Gly Pro Gly 200 205 Asp Trp Tyr Leu Lys Met Leu Gln Cys Cys Lys Cys Lys Gln Trp

PCT/US00/02237

				215					220					225
Phe	His	Glu	Ala	Cys 230	Val	Gln	Cys	Leu	Gln 235		Pro	Met	Leu	Phe 240
Gly	Asp	Arg	Phe	Tyr 245	Thr	Phe	Ile	Cys	Ser 250		. Cys	Ser	Ser	Gly 255
Pro	Glu	Tyr	Leu	Lys 260	Arg	Leu	Pro	Leu		Trp	Val	Asp	Ile	Ala 270
His	Leu	Cys	Leu		Asn	Leu	Ser	Val		His	Lys	Lys	Lys	Tyr 285
Phe	Asp	Ser	Glu		Glu	Leu	Met	Thr	Tyr 295		Asn	Glu	Asn	Trp
Asp	Arg	Leu	His	Pro 305	Gly	Glu	Leu	Ala	Asp 310		Pro	Lys	Ser	Glu 315
Arg	Tyr	Glu	His	Val 320	Leu	Glu	Ala	Leu	Asn 325		Tyr	Lys	Thr	Met 330
Phe	Met	Ser	Gly	Lys 335	Glu	Ile	Lys	Lys	Lys 340	Lys	His	Leu	Phe	Gly 345
Leu	Arg	Ile	Arg	Val 350	Pro	Pro	Val	Pro	Pro	Asn	Val	Ala	Phe	Lys 360
Ala	Glu	Lys	Glu	Pro 365	Glu	Gly	Thr	Ser	His 370	Glu	Phe	Lys	Ile	Lys 375
Gly	Arg	Lys	Ala	Ser 380	Lys	Pro	Ile	Ser	Asp 385	Ser	Arg	Glu	Val	Ser
Asn	Gly	Ile	Glu	Lys 395	Lys	Lys	Lys	Lys	Lys 400	Ser	Val	Gly	Arg	Pro 405
Pro	Gly	Pro	Tyr	Thr 410	Arg	Lys	Met	Ile	Gln 415	Lys	Thr	Ala	Glu	Pro 420
Leu	Leu	Asp	Lys	Glu 425	Ser	Ile	Ser	Glu	Asn 430	Pro	Thr	Leu	Asp	Leu 435
	_			440					Thr 445					450
				455					Ser 460					465
Thr	Ser	Ser	Ser	11e 470	Ser	Arg	His	Tyr	Gly 475	Leu	ser	Asp	Ser	Arg 480
_	_			485	_				Pro 490					495
	_		_	500					Arg 505					510
				515		-	_		Glu 520					525
		-		530					Leu 535					540
				545					Asn 550					Tyr 555
	-			560	-			-	Gly 565		_		_	570
				575				Gly	Lys 580	Val	Gln	Tyr		Val 585
Glu	Trp	Glu	Gly	Ala 590	Thr	Ala	Ser							

<210> 3

<211> 534

<212> PRT

<213> Homo sapiens

<220>

<400> 3

<221> misc-feature <223> Incyte ID No.: 116208CD1

Met Ara Ala Leu His Leu Leu Lys Ser Gly Cys Ser Pro Ala Val 1 10 Lys Ile Arg Glu Leu Tyr Arg Arg Arg Tyr Pro Arg Thr 20 25 Leu Glu Gly Leu Ser Ile Lvs Ser Ser Val Pho 35 40 45 Ser Leu Asp Gly Gly Ser Ser Pro Val Glu Pro Asp Leu Ala Val 50 55 Ile His Ser Leu Pro Ser Thr Ser Val Thr Pro His Ser 65 70 75 Pro Ser Ser Pro Val Gly Ser Val Leu Leu Gln Asp Thr Lys Pro 80 85 90 Thr Phe Glu Met Gln Gln Pro Ser Pro Pro Ile Pro Pro Val His 95 100 105 Pro Asp Val Gln Leu Lys Asn Leu Pro Phe Tyr Asp Val Leu Asp 110 115 Val Leu Ile Lys Pro Thr Ser Leu Val Gln 125 130 135 Phe Gln Glu Lys Phe Phe Ile Phe Ala Leu Val 140 145 150 Arg Glu Ile Cys Ile Ser Arg Asp Phe Leu Pro Gly Gly Arg Arg 155 160 165 Asp Tvr Thr Val Gln Val Gln Leu Arg Leu Cvs Leu Ala Glu Thr 170 175 Ser Cys Pro Gln Glu Asp Asn Tyr Pro Asn Lys 185 190 195 Val Asn Gly Lys Leu Pro Leu Pro Gly 200 205 210 Asn Gly Ile Glu Gln Lys Arg Pro Gly Arg Pro Leu Asn Ile 215 220 225 Ser Leu Val Arg Leu Ser Ser Ala Val Pro Asn Gln Ile Ser 230 235 240 Ser Trp Ala Ser Glu Ile Gly Lys Asn Tvr Ser Met Ser Val 245 250 255 Leu Val Arg Gln Leu Thr Ser Ala Met Leu Leu Gln Arg Leu 260 265 270 Met Lys Gly Ile Arg Asn Pro Asp His Ser Arg Ala Leu Ser 275 280 285 Glu Lys Leu Thr Ala Asp Pro Asp Ser Glu Ile Ala Thr Thr 290 295 Leu Arg Val Ser Leu Met Cys Pro Leu Gly Lys Met Arg Leu 305 310 315 Thr Ile Pro Cys Arg Thr 320 325 330 Asp Ala Ala Leu Tvr Leu Gln Met Asn Glu Lys Lys Pro Thr Trp 335 340 Ile Cys Pro Val Cys Asp Lys Lys Ala Ala Tyr Glu Ser Leu Ile 350 355 360 Leu Asp Gly Leu Phe Met Glu Ile Leu Asn Asp Cys Ser Asp Val 365 370 375 Glu Ile Lys Phe Gln Glu Asp Gly Ser Trp Cys Pro Met Arg 380 385 390 Lys Lys Glu Ala Met Lys Val Ser Ser Gln Pro Cys Thr Lys 395 400 405 Ile Glu Ser Ser Ser Val Leu Ser Lys Pro Cys Ser Val Thr Val 410 415 420 Ala Ser Glu Ala Ser Lys Lys Lys Val Asp Val Ile Asp Leu Thr 425 430

PCT/US00/02237

```
Ile Glu Ser Ser Ser Asp Glu Glu Glu Asp Pro Pro Ala Lys Arg
                440
                                     445
                                                          450
Lvs Cvs Ile Phe Met Ser Glu Thr Gln Ser Ser Pro Thr Lvs Glv
                                     460
                                                          465
                                                          Thr
Val Leu Met Tyr Gln Pro Ser Ser Val Arg
                470
                                     475
Ser Val Asp Pro Ala Ala Ile Pro Pro Ser Leu Thr Asp Tyr
                                                          Ser
                485
                                     490
                                                          495
Val Pro Phe His
               His Thr Pro Ile Ser Ser Met Ser Ser Asp
                                                          Leu
                500
                                     505
                                                          510
Pro Gly Glu Gln Arg Arg Asn Asp Ile Asn Asn Glu Leu Lys
                                                          Len
                515
                                     520
Gly Thr Ser Ser Asp Thr Val Gln Gln
                530
```

```
<210> 4
<211> 255
<212> PRT
<213> Homo sapiens
```

<400>

<221> misc feature <223> Incyte ID No.: 179261CD1

Met Lvs Ile Ile Ser Glu Asn Val Pro Ser Tyr Lys Thr His Glu 10 15 Ser Leu Thr Leu Pro Arg Arg Thr His Asp Ser Glu Lys Pro Tyr 20 25 ั ร ก Glu Tyr Lys Glu Tyr Glu Lys Val Phe Ser Cys Asp Leu Glu Phe 35 40 45 Asp Glu Tyr Gln Lys Ile His Thr Gly Gly Lys Asn Tyr Glu Cys 50 55 60 Asn Gln Cys Trp Lys Thr Phe Glv Ile Asp Asn Ser Ser Met Leu 70 65 Gln Leu Asn Ile His Thr Gly Val Lys Pro Cys Lys Tyr Met Glu 80 85 9 0 Tyr Gly Asn Thr Cys Ser Phe Tyr Lys Asp Phe Asn Val Tvr Gln 95 100 105 Ile His Asn Glu Lys Phe Tyr Lys Cys Lys Glu Tyr Arg Arg 110 115 120 Phe Glu Arg Val Gly Lys Val Thr Pro His 125 130 135 Asp Gly Glu Lys His Phe Glu Cys Ser Phe Cys Gly Lys Ser Phe 150 Arg Val His Ala Gln Leu Thr Arg His Gln Lys Ile His Thr Asp 155 160 165 Glu Lys Thr Tyr Lys Cys Met Glu Cys Gly Lys Asp Phe Arg Phe 170 175 180 His Ser Gln Leu Thr Glu His Gln Arg Ile His Thr Gly Glu Lys 185 190 195 Pro Tyr Lys Cys Met His Cys Glu Lys Val Phe Arg Ile Ser Ser 200 205 210 Gln Leu Ile Glu His Gln Arg Ile His Thr Gly Glu Lys Pro Tyr 215 220 225 Ala Cys Lys Glu Cys Gly Lys Ala Phe Gly Val Cys Arg Glu Leu 230 235 Ala Arg His Gln Arg Ile His Thr Gly Lys Tyr Cys Gly Trp Ile 245 250

<220>

PCT/US00/02237

<210> 5 <211> 562 <212> PRT <213> Homo sapiens <220> <221> misc feature <223> Incyte ID No.: 259161CD1 <400> 5 Met Ala Ser Val Ser Ala Leu Thr Glu Glu Leu Asp Ser Ile Thr 15 Ala Val Glu Ile Gln Ile Gln Glu Leu Thr 20 25 30 Arg Gln Gln Glu Leu Ile Gln Lys Lys Lys Val Leu Thr Lys Lys 35 40 45 Ile Lys Gln Cys Leu Glu Asp Ser Asp Ala Gly Ala Ser Asn Glu 50 55 Ser Ser Pro Ala Ala Trp Asn Lys Glu Asp Trp 65 Ser Gly Lys Val Lys Asp Ile Leu Gln Asn Val Phe Lys 80 85 90 Lvs Phe Arg Pro Leu Gln Leu Glu Thr Ile Asn Val Thr Met Ala 95 100 105 Leu Val Met Pro Thr Gly Gly Lys Gly Lys Glu Val Phe Ser 110 115 Leu Cys Tyr Gln Leu Ala Leu Cys Ser Leu 125 130 135 Ile Cys Pro Leu Ile Ser Leu Met Glu Asp Gln Leu Met Val 150 140 145 Leu Lys Gln Leu Gly Ile Ser Ala Thr Met 155 160 165 Lys Trp Val His Ala Glu Met Val Asn Lys Ser Lys Glu His Val 170 175 Asn Ser Glu Leu Lys Leu Ile Tyr Val Thr Pro Glu Lys Ile Ala 185 190 Ser Lys Met Phe Met Ser Arg Leu Glu Lys Ala Tyr Glu Ala 205 200 210 Arg Arg Phe Thr Arg Ile Ala Val Asp Glu Val His Cys Cys ser 215 220 Gln Trp Gly His Asp Phe Arg Pro Asp Tyr Lys Ala Leu Gly Tle 230 235 Leu Lys Arg Gln Phe Pro Asn Ala Ser Leu Ile Gly Leu Thr Ala 255 245 250 Thr Ala Thr Asn His Val Leu Thr Asp Ala Gln Lys Ile Leu Cys 260 265 Ile Glu Lys Cys Phe Thr Phe Thr Ala Ser Phe Asn Lys Asp 275 280 285 Val Arg Phe Val Ile His His Ser Met Ser Lvs Ser Met Glu Asn 290 295 300 Tyr Tyr Gln Glu Ser Gly Arg Ala Gly Arg Asp Asp Met Lys Ala 305 310 315 Asp Cys Ile Leu Tyr Tyr Gly Phe Gly Asp Ser Ile Phe Arg Ile 320 325 330 Ser Met Val Val Met Glu Asn Val Glv Gln Gln Lvs Leu Tvr Glu 345 335 340 Met Val Ser Tvr Cys Gln Asn Ile Ser Lys Cys Arg Arg Val Leu 350 355 360 Met Ala Gln His Phe Asp Glu Val Trp Asn Ser Glu Ala Cys Asn

PCT/US00/02237

```
370
                                                           375
Lys Met Cys Asp Asn Cys Cys Lys Asp Ser Ala Phe Glu Arg Lys
                380
                                                           390
                                     385
                    Cys Arg Asp Leu Ile Lys Ile Leu Lys Gln
Asn Ile Thr Glu
                Tvr
                395
                                     400
Ala Glu Glu Leu Asn Glu Lys Leu Thr Pro
                                         Leu Lys Leu Ile Asp
                                     415
                                                           420
Ser Trp Met Gly Lys Gly Ala Ala Lys Leu
                                         Arg Val Ala Glv Val
                425
                                     430
                                                          435
                                                          Ala
Val Ala Pro Thr Leu
                    Pro Arg Glu Asp Leu
                                                          450
                440
                                     445
His Phe Leu Ile Gln Gln Tyr Leu Lys Glu
                                         Asp Tyr Ser Phe
                455
                                     460
                                                          465
                                         Gly Pro Lys Ala
                                                          Asn
Ala Tyr Ala Thr
                Ile
                    Ser Tyr Leu Lys
                                    Tlo
                470
                                     475
Leu Leu Asn Asn Glu Ala His Ala Ile Thr Met Gln Val Thr Lys
                                                          495
                485
                                     490
Ser Thr Gln Asn Ser Phe Arg Ala Glu Ser
                500
                                     505
                                                          510
Ser Glu Gln Gly Asp Lys Lys Ile Gly Gly Lys Lys Phe Gln
                                                         Ala
                                     520
                                                          525
                515
Thr Ser Arg Arg Leu Gln Thr Cys
                                    Phe Ser Asn Leu Val
                                                          Leu
                530
                                     535
                                                          540
Arg Ile Gln Glu Leu Arg Lys Glu Lys Ser Met Met Pro Asp Met
                                     550
                                                          555
                545
Thr Val Thr Lys Phe Ser Asn
                560
```

```
<210> 6
<211> 432
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
```

<223> Incyte ID No.: 320087CD1

<400> 6 Met Ser Glu Leu Lys Asp Cys Pro Leu Gln Phe His Asp Phe Lys 10 15 Asp His Leu Lys Val Cys Pro Arg 20 25 3.0 Ala Arg Ser Glu Asp Asp Gly Ile Gly Ile Glu Glu Leu Asp Thr 35 40 45 Leu Gln Leu Glu Leu Glu Thr Leu Leu Ser Ser Ala Ser Arg Arg 55 60 50 Leu Arg Val Leu Glu Ala Glu Thr Gln Ile 65 70 Asp Lys Lys Gly Asp Arg Arg Phe Leu Lys Leu Gly Arg Asp His 80 85 90 Glu Leu Gly Ala Pro Pro Lys His Gly Lys Gln Lys 95 100 105 Leu Glu Gly Lys Ala Gly His Gly Pro Gly Pro Gly Pro Gly Arg 110 115 120 Pro Lys Ser Lys Asn Leu Gln Pro Lys Ile Gln Glu Tyr Glu Phe 125 130 Thr Asp Asp Pro Ile Asp Val Pro Arg Ile Pro Lys Asn Asp Ala 150 140 145

PCT/US00/02237

```
Asn Arg Phe Trp Ala Ser Val Glu Pro Tyr Cys Ala Asp Ile
                                                           165
                                      160
                155
                                                          Pro
   Ser Glu Glu Val Arg Thr Leu Glu Glu Leu Leu Lys Pro
                 170
                                      175
                                                           180
                                 Ile Pro
                                          Pro Leu Gly Lys
                                                          Hic
    Asp Glu Ala Glu His Tvr Lvs
                185
                                      190
                                                           195
    Ser Gln Arg Trp
                    Ala Gln Glu Asp Leu Leu Glu Glu Gln Lys
                200
                                      205
                                                           210
   Gly Ala Arg Ala
                    Ala Ala Val Ala Asp
                                          LVS LVS LVS GlV
                                                           Leu
                                                           225
                 215
                                      220
   Gly Pro Leu Thr Glu Leu Asp Thr
                                                          LOU
                                     Lys Asp Val Asp Ala
                230
                                      235
                                                           240
   Lys Lys Ser Glu
                                      Gln
                                          Pro Glu Asp Gly
                                                          Cvs
                                                           255
                245
                                      250
   Phe Gly Ala Leu Thr Gln Arg Leu Leu Gln Ala Leu Val Glu
                 260
                                      265
                                                           270
Glu Asn Ile Ile Ser Pro Met Glu Asp Ser Pro Ile Pro Asp Met
                275
                                      280
                                                           285
   Gly Lys Glu Ser Gly Ala Asp Gly Ala Ser Thr Ser Pro Arg
                290
                                      295
                                                           300
Asn Gln Asn Lvs Pro
                    Phe Ser Val Pro
                                          Thr Lys Ser Leu
                                                          Glu
                                      His
                                                          315
                305
                                      310
Ser Arg Ile Lys Glu Glu Leu Ile Ala Gln Gly Leu Leu Glu
                                                          Ser
                320
                                      325
                                                          330
   Asp Arg Pro Ala Glu Asp Ser Glu Asp Glu Val Leu Ala
                                                          Glu
                335
                                      340
                                                           345
    Arg Lys Arg Gln
                    Ala Glu Leu Lys
                                      Ala Leu Ser Ala His
                                                          Asn
                350
                                      355
                                                          360
                    Asp Leu Leu Arg Leu Ala Lys Glu Glu
   Thr Lys Lys His
                                                          Val
                365
                                      370
                                                          375
                    Arg Gln Arg Val Arg Met Ala Asp Asn
                                                          Glu
   Arg Gln Glu Leu
                380
                                      385
                                                          390
Val Met Asp Ala Phe Arg Lys Ile Met Ala Ala Arg Gln Lys
                                                          Lvs
                395
                                      400
                                                          405
                    Lys Glu Lys Asp Gln Ala Trp Lys Thr Leu
Arg Thr Pro Thr Lys
                410
                                      415
                                                          420
Lvs Glu Arg Glu Ser Ile Leu Lvs Leu Leu Asp Gly
                                      430
                425
```

```
<210> 7
<211> 799
<212> PRT
```

<220>

-400× 7

Met Pro Ser Gln Asn Tyr Asp Leu Pro Gln Lys Lys Gln Glu Lys 15 1 10 Phe Lys Asp Val Ala Val Met Thr Lys Phe Gln Glu Ala Val Thr 20 25 30 Asp Leu Thr Gln Arg Phe Ser Arg Glu Glu Leu Arg Leu Leu 35 40 45 Val Met Val Glu Asn Phe Lys Asn Leu Val Lys Leu Tyr Arg Asp 50 55 60 Ala Val Gly His Leu Pro Phe Gln Pro Asp Met Val Ser Gln Leu 70 65

<213> Homo sapiens

<221> misc-feature <223> 491271CD1

PCT/US00/02237

Glu	Ala	Glu	Glu	Lys 80	Leu	Trp	Met	Met	Glu 85		Glu	Thi	Glr	Arg 90
Ser	Ser	Lys	His	Gln 95	Asn	Lys	Met	Glu	Thr 100		Glr	Lys	Phe	Ala 105
	-	Tyr		110	Asn	Gln	Glu	Leu	Ser 115	Cys	Trp	Glr	ı Ile	120
Lys	Gln	Val	Ala	Ser 125	Glu	Leu	Thr	Arg	Cys 130	Leu	Gln	Gly	Lys	Ser 135
Ser	Gln	Leu	Leu	Gln 140	Gly	Asp	Ser	Ile	Gln 145	Val	Ser	Glu	Asn	Glu 150
Asn	Asn	Ile	Met	Asn 155	Pro	Lys	Gly	Asp	Ser 160	Pro	Ile	Tyr	Ile	Glu 165
		Glu		170		Trp			175				Gly	180
		Leu		185		Gln			190		Gly		Gln	195
		Lys		200		Gln		_	205					210
				215		Ile	-		220					225
-	_	Gly		230	-	Gly			235	Ser			Ser	240
	-	Leu		245		Glu			250		Cys		Glu	255
		Gly		260	-		Pro		265				Pro	270
		Thr		275		Cys			280				Leu	285
		Gln		290	His	Asn	Gly		295			Arg	-	300
		Gly His	_	305	Glu				310	Cys			Cys	315
		Phe		320		-	Gly	_	325	Ile		Tyr	Arg	330
-		Gly		335					340				Lys	345
		Gln		350		Phe	_	_	355					360 Thr
				365		Cys			370					375 Gly
				380 Leu					385				Gly	390
-	Pro			395	_	Glu			400	Gly	٠.		Gln	405
- Ala	His	Phe	-	410		Gln			415	Thr	Gly	Glu	Lys	420 Pro
Tyr	Lys	Cys		425 Val			Lys		430	Ser	His	Asn	Ser	435 Pro
				440		Val	His	Thr	445 Gly	Glu	Lys	Pro	Tyr	450 Lys
				455					460					465
Cys	Glu	Ala	Cys	Gly 470	Lys	Gly	Phe	Thr	Arg 475	Asn	Thr	Asp	Leu	His 480
Ile	His	Phe	Arg	Val 485	His	Thr	Gly	Glu	Lys 490	Pro	Tyr	Lys	Суѕ	Lys 495
Glu	Cys	_	-	500		Ser			505		Leu	Gln	Val	His 510
	Asn			515		Glu			520		Cys			Cys 525
GJĀ	Lys	Gly	Phe	ser	Gln	Ser	Ser	Lys	Leu	Gln	Thr	His	Gln	Arg

10/91

"我看帮你要你看她就是重新的一颗脚上的一脚就看得一切的人的心,你就想到了这个人的人的大哥

PCT/US00/02237

```
530
                                      535
                                                            540
Val His Thr Gly Glu Lys Pro Tyr Arg Cys Asp Val Cys Gly
                                                           Lys
                 545
                                      550
    Phe Ser Tyr Ser Ser Asn Leu Lys Leu His Gln Val
                                                           His
                 560
                                      565
                                                            570
Thr Gly Glu Lys Pro Tyr Lys Cys Glu Glu Cys Gly Lys Gly
                                                            Phe
                 575
                                      580
                                                            585
    Trp Arg Ser Asn Leu His Ala His Gln Arg Val His Ser Glv
                 590
                                      595
                                                            600
Glu Lys Pro Tyr Lys
                    Cys Glu Gln Cys Asp
                                                           Gln
                 605
                                      610
                                                            615
Ala Ile Asp Phe Arg
                    Val His Gln Arg Val
                                                           Lys
                 620
                                      625
                                                            630
    Tyr Lys Cys Gly
                     Val Cys Gly Lys Gly
                                          Phe Ser Gln Ser
                                                           Ser
                 635
                                      640
Gly Leu Gln Ser His Gln Arg Val His Thr
                                          Gly Glu Lys Pro
                                                           Tyr
                 650
                                      655
                                                            660
    Cys Asp Val Cys Gly Lys Gly Phe Arg
                                              Ser Ser Gln Phe
                 665
                                      670
                                                           675
    Tyr His Gln Arg Gly His Thr Gly
                                     Glu
                                         Lvs Pro Tyr Lys
                                                           Cvs
                680
                                      685
                                                           690
   Glu Cys Gly Lys Gly Phe Gly Arg
                                     Ser
                                          Leu Asn Leu Arg
                                                           His
                695
                                      700
                                                           705
His Gln Arg Val His Thr Gly Glu Lys
                                     Pro
                                         His Ile Cvs Glu Glu
                710
                                      715
                                                           720
   Gly Lys Ala Phe Ser Leu Pro Ser
                                     Asn
                                                           Leu
                725
                                      730
                                                           735
Gly Val His Thr Arg Glu Lys Leu Phe
                                     Lvs
                                                           Glv
                740
                                      745
                                                           750
Lys Gly Phe Ser Gln Ser Ala Arg Leu
                                     Glu Ala His Gln Arg
                                                           Va1
                755
                                     760
                                                           765
   Thr Gly Glu Lys
                    Pro Tyr Lys Cys
                                     Asp
                                         Ile Cys Asp Lys
                                                           Asp
                770
                                     775
Phe Arg His Arg Ser Arg Leu Thr Tyr His Gln Lys Val His Thr
                785
                                     790
                                                           795
Gly Lys Lys Leu
```

```
<210> 8
<211> 137
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 585172CD1
```

<400> 8 Met Leu Ser Gly Arg Leu Val Leu Gly Leu Val Ser Met Ala Gly 10 Arg Val Cys Leu Cys Gln Gly Ser Ala Gly Ser Gly Ala Ile Gly 20 25 30 Pro Val Glu Ala Ala Ile Arg Thr Lys Leu Glu Glu Ala Leu Ser 35 40 Pro Glu Val Leu Glu Leu Arg Asn Glu Ser Gly Gly His Ala Val 50 60

Pro Pro Gly Ser Glu Thr His Phe Arg Val Ala Val Val Ser Ser 65 70 75 Arg Phe Glu Gly Leu Ser Pro Leu Gln Arg His Arg Leu Val His

PCT/US00/02237

```
Ala Ala Leu Ala Glu Glu Leu Gly Gly Pro Val His Ala Leu Ala
                  95
                                      100
                                                          105
Ile Gln Ala Arg Thr Pro Ala Gln Trp Arg Glu Asn Ser Gln Leu
                 110
                                      115
                                                          120
Asp Thr Ser Pro Pro Cys Leu Gly Gly Asn Lys Lys Thr Leu Gly
                 125
                                     130
Thr Pro
<210> 9
<211> 230
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 615200CD1
<400> 9
```

85

Met Val Gly Ala Gly Ile Ser Thr Pro Ser Gly Ile Pro Asp Phe 10 Arg Ser Pro Gly Ser Gly Leu Tyr Ser Asn Leu Gln Gln Tyr Asp 25 30 Leu Pro Tyr Pro Glu Ala Ile Phe Glu Leu Pro Phe Phe His 35 40 45 Lys Pro Phe Phe Thr Leu Ala Lys 50 55 60 Asn Tyr Lys Pro Asn Ile Thr His Tyr Phe Leu Arg Leu Leu His 65 70 75 Asp Lys Gly Leu Leu Leu Arg Leu Tyr Thr Gln Asn Ile Asp Gly 80 85 90 Leu Glu Arg Val Ser Gly Ile Pro Ala Ser Lys Leu Val Glu Ala 95 100 105 His Gly Thr Phe Ala Ser Ala Thr Cvs Thr Val Cvs Gln Arg Pro 110 115 120 Phe Pro Gly Glu Asp Ile Arg Ala Asp Val Met Ala Asp Arg Val 125 130 135 Pro Arg Cys Pro Val Cys Thr Gly Val Val Lys Pro Asp Ile Va1 140 145 150 Phe Gly Glu Pro Leu Pro Gln Arg Phe Leu Leu His Val Val 155 160 165 Pro Met Ala Asp Leu Leu Leu Ile Leu Gly Thr Ser Leu 170 175 180 Glu Val Glu Pro Phe Ala Ser Leu Thr Glu Ala Val Arg Thr Gln 185 190 Phe Pro Asp Cys Ser Ser Thr Gly Thr Trp Trp Gly Pro Trp Leu 205 200 Gly Ile Leu Ala Ala Gly Thr Trp Pro Ser Trp Gly Thr Trp Phe 220 215 Thr Ala Trp Lys Ala 230

```
<210> 10
<211> 446
<212> PRT
<213> Homo sapiens
```

<221> misc-feature

<220>

PCT/US00/02237

<223> Incvte ID No.: 997067cD1 Met Glu Thr Gln Ala Asp Leu Val Ser Gln Glu Pro Gln Ala Leu 15 1 1.0 Leu Asp Ser Ala Leu Pro Ser Lys Val Pro Ala Phe Ser Asp Lvs 20 25 30 Asp Ser Leu Gly Asp Glu Met Leu Ala Ala Ala Leu Leu Lys Ala 35 40 45 Lys Ser Gln Glu Leu Val Thr Phe Glu Asp Val Ala Val Tyr Phe 55 50 60 Ile Arg Lys Glu Trp Lys Arg Leu Glu Pro Ala Gln Arg Asp Leu 70 75 65 Tyr Arg Asp Val Met Leu Glu Asn Tyr Gly Asn Val Phe Ser Leu 90 80 85 Asp Arg Glu Thr Arg Thr Glu Asn Asp Gln Glu Ile Ser Glu Asp 95 100 105 Thr Arg Ser His Glv Val Leu Leu Gly Arg Phe Gln Lvs Asp Ile 110 115 120 Phe Lys Glu Ala Tyr Glu Arg Glu Val Ser Ser Gln Gly Leu Lys 130 135 125 Gly Asn Ser Pro Gly Leu Lys Arg Pro Leu Glu Arg Leu Asn Arg 140 145 150 Lvs Met Pro Asp Phe Gly Gln Val Thr Val Glu Glu Lys Thr 160 155 Ser Pro Arg Gly Glu Arg ser Glu Lys Tyr Asn Asp Phe 170 175 180 Phe Thr Val Asn Ser Asn Leu Ile Ser His Gln Arg Leu Pro Val 185 190 195 Gly Asp Arg Pro His Lys Cys Asp Glu Cys Ser Lys Ser Phe Asn 205 210 200 Ile Gln His Gln Arg Ile His Thr Gly Glu Arg Thr Ser Asp Leu 215 220 225 Lys Pro Tyr Glu Cys Asn Glu Cys Gly Lys Ala Phe Ser 230 235 240 Ser His Leu Ile Gln His Gln Arg Ile His Thr Gly Glu Lys Pro 250 255 245 Ala Tvr Glu Cvs Ser Asp Cvs Glv Lvs Thr Phe Ser Cys Ser Ser 265 270 260 Leu Ile Leu His Arg Arg Ile His Thr Gly Glu Lys Glu 280 285 275 Asn Glu Cys Gly Lys Thr Phe Ser Trp Ser Ser Thr 290 295 300 His Thr Gly Glu Lys Pro Tyr Ala Cys Asn His His Gln Arg Ile 305 310 315 Thr Leu Ile His Glu Cys Gly Lys Ala Phe Ser Arg Ser Ser His 320 325 330 Cys Gln Arg Ile His Thr Gly Glu Lys Pro Tyr 335 340 345 Gly Lys Ala Phe Ser Gln Ser Ser His Leu Tyr Gln His Gln Ara 350 355 360 Ile His Thr Gly Glu Lys Pro Tyr Glu Cys Met Glu Cys Gly Gly 370 375 365 Lys Phe Thr Tyr Ser Ser Gly Leu Ile Gln His Gln Arg Ile His 380 385 390 Thr Gly Glu Asn Pro Tyr Glu Cys Ser Glu Cys Gly Lys Ala Phe 395 400 405 Leu Val Arg His Gln Arg Ile His Tyr Ser Ser Ala Thr Gly 420 415 410

Glu Lys Pro Leu Asn Gly Ile Gly Met Ser Lys Ser Ser Leu Arg

435

WO 00/44900

PCT/US00/02237

```
425 430
Val Thr Thr Glu Leu Asn Ile Arg Glu Ser Thr
440 445
```

<210> 11 <211> 428 <212> PRT <213> Homo sapiens

<220> <221> misc-feature

<400> 11

<223> Incyte ID NO.: 1443262CD1

Met Glu Pro Leu Lys Val Glu Lys Phe Ala Thr Ala Asn Arg Gly 5 10 15 Leu Asn Gly Leu Arg Ala Val Thr Pro Leu Arg 30 20 25 Ara Pro Thr Val Phe Arg Ser Asp 45 35 40 Arg Cys Leu Leu Gly Lys Glu Lys Leu Met Asp 60 50 55 Cys Ser Ala Lvs Cvs Arg Cvs Ser Gln Cvs Arg Val Ala Lys Tyr 70 75 65 Len Pro Asp His Lys Arg Glu Cys Lys Cys Gln Lys Lys Ala Trp 85 90 80 T.011 Arg Tyr Pro Pro Asp Ser Val Arg Leu Lys Ser Cys Lys Pro 105 95 100 Leu Met Asp Gly Ser Val Val Phe 120 115 110 Tyr Asp Leu Glu Ser Asn Ile Asn Glu Lys Leu Tyr Ser 130 135 125 Thr Lys Glu Gly Leu Arg Gln Leu Val Met Leu Thr Glu Asp Lys 150 140 145 Phe Gln His Phe Met Arg Glu Glu Ile Gln Asp Ala Ser Gln Leu 165 160 155 Pro Ala Phe Asp Leu Phe Glu Ala Phe Ala Lys Val Ile 170 175 180 Ile Cys Asn Ala Glu Met Gln Glu Val Gly Asn Ser Phe Thr 195 185 190 Pro Ile Ser Leu Leu Asn His Ser Cys Asp Ser Gly Leu Tyr Pro 200 205 Phe Asn Gly Pro His Leu Leu Leu Arg Ala Asn Cvs Ser Ile Val 215 220 Val Arg Asp Ile Glu Val Gly Glu Glu Leu Thr Ile Cys Tyr 240 230 235 Asp Met Leu Met Thr Ser Glu Glu Arg Arg Asp 255 250 245 Gln Tyr Cys Phe Glu Cys Asp Cys Phe Arg Cys Gln Thr Gln ASD 260 265 Lys Asp Ala Asp Met Leu Thr Gly Asp Glu Gln Val Trp Lys 285 280 275 Val Gln Glu Ser Leu Lys Lys Ile Glu Glu Leu Lys Ala His Trp 300 290 295 Lys Trp Glu Gln Val Leu Ala Met Cys Gln Ala Ile Ile Ser Ser 315 305 310 Asn Ser Glu Arg Leu Pro Asp Ile Asn Ile Tyr Gln Leu Lys Val 330 325 320

PCT/US00/02237

```
Asp Cys Ala Met Asp Ala Cys Ile Asn Leu Gly Leu Leu Glu
                335
                                     340
                                                           345
        Leu Phe Tyr Gly Thr Arg Thr Met Glu Pro Tyr Arg
                                                          Tla
                350
                                     355
                                                           360
                Ser His Pro Val Arg Gly
    Phe Pro Glv
                                         Val Gln Val Met
                                                          Lvs
                                     370
                                                           375
                365
Val Gly Lys Leu Gln Leu His Gln Gly Met
                                         Phe Pro Gln Ala Met
                380
                                     385
   Asn Leu Arg Leu Ala Phe Asp Ile Met Arg Val
                                                 Thr His Gly
                395
                                     400
                                                          405
Arg Glu His Ser Leu Ile Glu Asp Leu Ile Leu Leu Leu Glu
                                                          Glu
                410
                                     415
                                                          420
Cys Asp Ala Asn Ile Arg Ala Ser
                425
```

```
<210> 12
<211> 590
<212> PRT
<213> Homo sapiens
```

<400> 12 Met Ala Glu Asp Trp Leu Asp Cys Pro Ala Leu Gly Pro Gly Trp 10 15 Va 1 Ser Gly Ala Thr Cys Gly Ara 20 25 30 Thr Gly ser Ser Asp Thr Tyr Tyr Gln Ser Pro 35 40 45 Lys Val Glu Leu Thr Arg Tyr Leu Gly Pro Thr 60 50 Lys Leu Phe Asp Phe Lys Gln Gly Ile Leu Cys Tvr Pro Ala Pro 70 75 65 Ala His Pro Val Ala Val Ala Ser Lys Lys Arg Lys Ser 85 90 Ωn Arg Pro Ala Lys Thr Arg Lys Arg Gln Val Gly Pro Gln Ser Glv 100 95 Glu Val Arg Lys Glu Ala Pro Arg Asp Glu Thr Lys Ala Asp Thr 115 120 110 Asp Thr Ala Pro Ala Ser Phe Pro Ala Pro Gly Cys Cys Glu Asn 125 130 Cys Gly Ile Ser Phe Ser Gly Asp Gly Thr Gln Arg Gln Arg 150 140 145 Asp Cvs Arg Ala Gln Arg Ile Ala Phe Asn Lys Thr Leu Cys Lys 155 160 165 Arg Glu Gln Arg Met Phe Lys Arg Val Gly Cys Gly Glu Cys Ala 175 170 Ala Cys Gln Val Thr Glu Asp Cys Gly Ala Leu 190 195 185 Leu Gln Leu Pro His Asp Val Ala Ser Gly Leu Phe Cvs Lvs CVS 200 205 210 Leu Arg Ile Val Glu Arg Ser Arg Gly Cys Glu Arg Arg Arg Cys 215 220 225 Cys Gln Thr Gln Glu Asp Cys Gly Val Cys Arg Gly Cys 230 235 240 Pro Ile Cys Leu Arg Pro Pro Arg Pro Gly Leu Arg Arg Gln Trp 245 250 255

<220> <221> misc-feature

<221> misc-reature <223> Incyte ID No.: 1521648CD1

PCT/US00/02237

```
Cys Val Gln Arg Arg Cys Leu Arg Gly Lys His Ala Arg Arg
                                       265
                 260
                                                            270
    Glv Glv Cvs
                 Asp
                     Ser Lys Met Ala Ala Arg Arg Arg Pro
                                                           GIV
                 275
                                       280
                                                            285
Ala Cla Pro Lou Pro
                     Pro Pro Pro Pro
                                      Ser Gln Ser Pro Glu
                                                           Pro
                 290
                                       295
                                                            300
Thr Glu Pro His
                 Pro
                     Arg Ala Leu Ala Pro
                                          Ser Pro Pro Ala Glu
                 305
                                       310
                                                            315
    Ile Tyr Tyr Cys
                         Asp Glu Asp Glu Leu Gln Pro Tyr
                                                           Thr
                 320
                                      325
                                                           330
Asn Arg Arg Gln Asn Arg Lys Cys Gly Ala Cys Ala Ala Cys
                                                           Leu
                 335
                                       340
                                                           345
Arg Arg Met Asp
                 Cys
                     Gly Arg Cys Asp Phe Cys Cys Asp Lys
                                                           Pro
                 350
                                      355
                                                           360
    Phe Gly Gly Ser Asn Gln Lys Arg Gln Lys Cys Arg Trp
                                                           Ara
                                      370
                 365
                                                           375
Gln Cys Leu Gln Phe Ala Met Lys Arg Leu Leu Pro Ser Val
                                                           Trp
                 380
                                      385
                                                           390
Ser Glu Ser Glu Asp Gly Ala Gly Ser Pro Pro Pro Tyr Arg Arg
                 395
                                      400
                                                           405
Arg Lys Arg Pro Ser Ser Ala Arg Arg His His Leu Gly Pro Thr
                 410
                                      415
                                                           420
Leu Lys Pro Thr Leu
                     Ala Thr Arg Thr Ala Gln Pro Asp His
                                                           Thr
                 425
                                      430
                                                           435
                     Gln Glu Ala Glv Glv
Gln Ala Pro Thr Lvs
                                          Gly Phe Val Leu
                 440
                                      445
                                                           450
                     Leu Val Phe Leu Arg Glu Gly Ala Ser
Pro Pro Gly Thr Asp
                                                          Ser
                 455
                                      460
                                                           465
Pro Val Gln Val Pro Gly Prc Val Ala Ala Ser Thr Glu Ala
                                                          Leu
                 470
                                      475
Leu Gln Val Lys Gln Glu Lys Ala Asp
                                      Thr Gln Asp Glu Trp
                                                           Thr
                 485
                                      490
                                                           495
Pro Gly Thr Ala
                 Val
                                      Val
                                                           Cys
                     Leu Thr Ser Pro
                 500
                                      505
                                                           510
Pro Ser Lys Ala
                 Val Asp Pro Gly Leu
                                     Pro
                                          Ser Val Lys Gln Glu
                 515
                                      520
                                                           525
Pro Pro Asp Pro
                 Glu Glu Asp Lys Glu Glu Asn Lys Asp Asp
                                                           Ser
                 530
                                      535
                                                           540
Ala Ser Lvs Leu
                Ala
                     Pro Glu Glu Glu Ala Gly Gly Ala Gly
                                                          Thr
                 545
                                      550
                                                           555
        Ile Thr Glu Ile Phe Ser Leu Gly Gly Thr Arg Phe
                                                          Arg
                 560
                                      565
                                                           570
                Trp
Asp Thr Ala Val
                    Leu Pro Arg Ser Lys Asp Leu Lys Lys
                                                          Pro
                 575
                                      580
                                                           585
                Gln
Gly Ala Arg Lys
                 590
```

<210> 13

<211> 479

<212> PRT <213> Homo sapiens

<220>

<221> misc-feature

<223> Incyte ID No.: 1685494CD1

<400> 13

Met Ala Thr Ala Leu Val Ser Ala His Ser Leu Ala Pro Leu Ser

PCT/US00/02237

1				5	5				1	0				15
Leu	Lys	Lys	Glu	Gly 20		a Arg	y Val	l Val	Arc 2	g Glu	ı Ası	His	s Ty	r Set
Thr	Trp	Glu	Gln	Gly 35		Lys	Leu	ı Glr		/ Asr	ı Ser	Lys	Gly	
Gly	Gln	Glu	Pro	Leu 50		Lys	Glr	ı Phe	Arg 55	g Glr	ı Lev	Arg	у Туг	Glu 60
Glu	Thr	Thr	Gly	Pro 65		Glu	Ala	. Leu	Ser 70		, Leu	Arg	g Glu	Leu 75
Cys	Gln	Gln	Trp	Leu 80		Pro	Glu	Thr	His 89		Lys	Glu	Glr	
Leu	Glu	Leu	Leu	Val 95		Glu	Gln	Phe	100		Ile	Leu	Pro	Lys 105
Glu	Leu	Gln	Ala	Arg 110		Gln	Glu	His	His 115		Glu	Ser	Arg	Glu 120
Asp	Val	Val	Val	Val 125		Glu	Asp	Leu	Glr 130	Leu	Asp	Leu	Gly	Glu 135
Thr	Gly	Gln	Gln	Val 140		Pro	Asp	Gln	Pro 145		Lys	Gln	Lys	11e
Leu	Val	Glu	Glu	Met 155	Ala	Pro		_	160					165
	_	His		170				Lys	175	Glu	Lys	Glu	Lys	Gly 180
			-	185		Asn	_	-	190		Val			195
	-	-	_	200				Gly	205		Ser			210
		His		215					220	Arg			Ala	225
		Glu		230		Tyr	Lys	Cys	Ser 235		Arg		Gln	240
		Gln		245				Glu	250	Ala			His	255
_	-	Lys		260			_	Val	Cys 265	Gln				Leu 270
				275					280	Lys				285
		_	_	290		Phe			Ser 295	Ser	His		Val	300
	Gln	-		305		Gly			310			-	Asn	315
-	-	-		320					325	Leu				330
_		His		335			Pro		340	Cys			-	Gly 345
_		Phe		350					355	Arg			_	360
		Gln		365					370	Glu		Gly	-	Thr 375
Phe		Gln		380				His	385	Gln	-	Ile		Ser 390
		_		395		Cys			400	Gly				405
				410		Arg			415		His		-	420
_				425			_		430	Ala		_		435
Ser		Leu		440					445	Asn -			_	Pro 450
-	Gln			Glu 455					460	Arg			Ser	Gly 465
Leu	Phe	Gln	His	Gln	Arg	Tyr	His	His	Lys	Asp	Lys	Leu	Ala	

17/91

y day in the state of the first of the second of the secon

<210> 14

PCT/US00/02237

470

475

15 Leu Val Leu Asp Leu Asn Phe Leu Leu Val Ser Leu Ala Trp Leu Leu Ala Phe Val Tyr Asn 40 45 35 Leu Pro His Thr Val Leu Thr Ser Leu Leu His Leu Gly Arg Gly 55 60 50 Val Leu Leu Ser Leu Leu Ala Leu Ile Glu 75 65 70 Thr Cys Gly Gly Leu Gln Ala Leu Cys Thr Leu Leu Tyr Ser CVS 80 85 ٩0 Gly His Leu Ala Ser Cvs Ser Gly Leu Glu Ser Leu Lys Leu Leu 100 105 95 His Gly Ala Leu Arg Ser Arg Glu Ile Leu His Arg Gly Val Leu 120 115 110 Asn Val Val Ser Ser Gly His Ala Leu Leu Arg Gln Ala Cys 130 135 125 Ala Met Ser Leu Val Ala Tyr Val Ile Asn Ile Cvs Ala Ile 140 145 150 Leu Ile Gly Thr Gln Asn Leu Phe Ser Len Cys 160 155 Val Leu Ala Leu Trp Asp Ala Val Thr Gly Pro Leu Trp Arg Met 175 170 Thr Asp Val Val Ala Ala Phe Leu Ala His 195 185 190 Leu Leu Trp Thr Pro Cys Gln Leu Ala Leu Val Ala Met Ala Ile 200 205 210 Ser Ala Ala Arg Leu Leu Ala Ser Phe Val Leu Glu Leu Leu Ala 215 220 Cys Val Leu Ala Val Val Asn Leu Thr Gly Leu Val Leu Leu Ala 240 230 235 Thr Val Thr Val Leu His Pro Asp Phe Thr Leu Arg Leu Ala 250 245 Gln Leu His Ala Arg Pro Ser Tyr His Arg Leu Gln Ala Leu Ser 265 260 Arg Glu Asp Val Met Arg Leu Ser Arg Leu Ala Leu Gly Ser Glu 285 275 280 Val Trp Ser Arg Ser Leu Gln Leu Ala Ser Trp Ala Trp Arg Arg 290 295 Ala Pro Gly Ala Pro Gln Gly Asp Pro Met Gly Pro Asn Arg Gly 310 305 Arg Val Phe Ser Val Arg Thr Arg Arg Gln Asp Thr Leu Pro Glu 330 320 325 Ala Gly Arg Arg Ser Glu Ala Glu Glu Glu Glu Ala Arg Thr 340 345 335 Arg Val Thr Pro Val Arg Gly Arg Glu Arg Leu Asn Glu Glu Glu

PCT/US00/02237

```
350
                                     355
                                                           360
Pro Pro Gly Gly Gln Asp Pro Trp Lys Leu Leu Lys Glu Gln Glu
                365
                                     370
                                                           375
Glu Arg Lys Lys Cys
                    Val Ile Cys Gln Asp Gln Ser Lys Thr
                                                          Val
                380
                                     385
                                                           390
Leu Leu Leu Pro Cys Arg His Leu Cys Leu Cys Gln Ala Cys
                                                          Thr
                395
                                     400
                                                           405
Glu Ile Leu Met Arg His Pro Val Tvr His Arg Asn Cvs Pro Leu
                410
                                     415
                                                           420
Cys Arg Arg Gly Ile Leu Gln Thr Leu Asn Val Tyr Leu
                                     430
```

<210> 15

<211> 320

<212> PRT

<213> Homo sapiens

<220>

<221> misc feature

<223> Incvte ID No.: 1864641CD1

Met Pro Lys Lys Thr Gly Ala Arg Lys Lys Ala Glu Asn Arg 10 Lys Gln Leu Arg Ala Ser Arg Ser Thr Ile Asp 25 30 20 Leu Ala Lys His Pro Cys Asn Ala Ser Met Glu Cys Asp Lys Cys 40 45 Gln Arg Arg Gln Lys Asn Arg Ala Phe Cys Tyr Phe Cys Asn Ser 50 55 60 Val Gln Lys Leu Pro Ile Cys Ala Gln Cys Gly Lys Thr Lys Cys 65 70 Met Met Lys Ser Ser Asp Cys Val Ile Lys His Ala Gly Tyr 85 80 Ser Thr Gly Leu Ala Met Val Gly Ala Ile Cys Asp Phe Cys Glu 95 100 105 Ala Trp Val Cys His Gly Arg Lys Cys Leu Ser Thr His Ala Cys 110 115 120 Ala Cys Pro Leu Thr Asp Ala Glu Cys Val Glu Cys Glu Arg Gly 130 135 Trp Asp His Gly Gly Arg Ile Phe Ser Cys Ser Phe Cys His 140 145 150 Asn Phe Leu Cys Glu Asp Asp Gln Phe Glu His Gln Ala Ser Cys 155 160 165 Gln Val Leu Glu Ala Glu Thr Phe Lys Cys Val Ser Cys Asn Arg 170 175 Leu Gly Gln His Ser Cys Leu Arg Cys Lys Ala Cys Phe Asp 190 195 185 Asp His Thr Arg Ser Lys Val Phe Lys Gln Glu Lys Gly Lys Gln 200 205 210 Pro Cys Pro Lys Cys Gly His Glu Thr Gln Glu Thr Lys Asp 215 220 225 Leu Ser Met Ser Thr Arg Ser Leu Lys Phe Gly Arg Gln Thr Gly 235 240 230 Gly Glu Glu Gly Asp Gly Ala Ser Gly Tyr Asp Ala Tyr Trp Lys 245 250 Asn Leu Ser Ser Asp Lys Tyr Gly Asp Thr Ser Tyr His Asp Glu 260 265 Glu Glu Asp Glu Tyr Glu Ala Glu Asp Asp Glu Glu Glu Glu Asp

<210> 16

PCT/US00/02237

```
<211> 179
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2444604CD1
```

<400> 16 Met Ala Ala Gly Phe Phe Gln Pro Phe Met Ser Pro Arg Phe Pro 10 Gly Gly Pro Arg Pro Thr Leu Arg Met Pro Ser Gln Pro Pro Ala 25 Cys Leu Pro Gly Ser Gln Pro Leu Leu Pro Gly Ala Met Glu Pro 35 40 Ser Pro Arg Ala Gln Gly His Pro Ser Met Gly Gly Pro Met Gln 50 55 60 Arg Val Thr Pro Pro Arg Gly Met Ala Ser Val Gly Pro Gln Ser 65 70 Tyr Gly Gly Gly Met Arg Pro Pro Pro Asn Ser Leu Ala Gly Pro 8 n 85 90 Gly Leu Pro Ala Met Asn Met Gly Pro Gly Val Arg Gly Pro Trp 95 100 105 Ala Ser Pro Ser Gly Asn Ser Ile Pro Tyr Ser Ser Ser 110 115 Gly Ser Tyr Thr Gly Pro Pro Gly Gly Gly Pro Pro Gly Thr 130 Pro Ile Met Pro Ser Pro Gly Asp Ser Thr Asn Ser Ser Glu Asn 140 145 Met Tyr Thr Ile Met Asn Pro Ile Gly Gln Gly Ala Gly Arg Ala 155 160 Asn Phe Pro Leu Gly Pro Gly Pro Glu Gly Pro Trp Pro Pro 170 175

20

PCT/US00/02237

Lуs	Ala	Lys	His		Lys	Суѕ	His	Ile	Cys 40	His	Lys	Lys	Leu	Tyr 45
Thr	Gly	Pro	Gly		Ala	Ile	His	Cys		Gln	Val	His	Lys	Glu 60
Thr	Ile	Asp	Ala		Pro	Asn	Ala	Ile		Gly	Arg	Thr	Asp	11e 75
Glu	Leu	Glu	Ile	Tyr 80	Gly	Met	Glu	Gly	Ile 85	Pro	Glu	Lys	Asp	Met 90
Asp	Glu	Arg	Arg		Leu	Leu	Glu	Gln	Lys 100	Thr	Gln	Glu	Ser	Gln 105
Lys	Lys	Lys	Gln	Gln 110	Asp	Asp	Ser	Asp	Glu 115	Tyr	Asp	Asp	Asp	Asp 120
ser	Ala		Ser	125					130	Pro			Pro	135
Gln				140					145				Pro	150
Pro	Gly			155					160				Met	165
Gly				170	Met				175				Pro	180
				185					190			Gln		Phe 195
-				200	Met				205				Pro	210
-		-	Ile	215					220				Gly	225
				230					235				Gln Ala	240
Gln				245	Pro	_			250			Lvs	Pro	255
				260	Ala				265	Val		-		270 Ser
			Ala	275		Glu			280				Lys	285
				290					295				Gly	300
Val				305	Lys				310				Thr	315
	-			320	Pro		Phe		325			Gln		330 Thr
			Thr	335		Thr			340				Pro	345 Ala
Ala				350	Lys				355				Ser	360 Ala
				365					370				Glu	375 Glu
				380					385				Arg	Pro
				395					400					405
				410					415				Gly	420
				425					430				Met	435
				440					445				Gln	450
				455					460				Gln -	465
Pro	Pro	Met	Val	Pro 470	Pro	Tyr	Gln	Gly	Gly 475	Pro	Pro	Arg	Pro	Pro 480

25

21/91

PCT/US00/02237

Met Gly Met Arg Pro Pro Val Met Ser Gln Gly Gly Arg Tyr 490

<210> 18 <211> 401 <212> PRT <213> Homo sapiens

<220>

<221> misc-feature <223> Incvte ID No.: 2572462CD1

<400> 18 Met Ala Ser Ser Pro Arg Pro Lys Met Asp Ala Ile Leu Thr Glu 10 Cys Phe Gln Lys Ser Gly Ala Ser Val Val Ala 20 25 30 Ile Ile Arg Lys Tyr Ile His Lys Tyr Pro Ser Leu Glu Leu Glu 35 40 45 Arg Arg Gly Tyr Leu Leu Lys Gln Ala Leu Lys Arg Glu Leu Asn 50 55 60 Lys Gln Val Lys Gly Lys Gly Ala Ser Gly Ser 70 65 75 Phe Val Val Val Gln Lys Ser Arg Lys Thr Pro Gln Lys Ser Arg 80 85 90 Asn Arg Lys Asn Arg Ser Ser Ala Val Asp Pro Glu Pro Gln Val 95 100 105 Lys Leu Glu Asp Val Leu Pro Leu Ala Phe 110 115 120 Pro Lys Glu Ala Ser Tyr Ser Leu Ile Arg Lvs Tvr Val Ser Gln 125 130 135 Tyr Tyr Pro Lys Leu Arg Val Asp Ile Arg Pro Gln Leu Leu Lys 140 145 Asn Ala Leu Gln Arg Ala Val Glu Arg Gly Gln Leu Glu Gln Ile 155 160 165 Thr Gly Lys Gly Ala Ser Gly Thr Phe Gln Leu Lys Lys Ser Gly 170 175 180 Lys Pro Leu Leu Gly Gly Ser Leu Met Glu Tyr Ala Ile 185 190 195 Ser Ala Ile Ala Ala Met Asn Glu Pro Lys Thr Cys Ser Thr Thr 200 205 210 Ala Leu Lys Lys Tyr Val Leu Glu Asn His ser 215 220 225 Asn Tvr Gln Met His Leu Leu Lys Lys Thr Glu 230 235 240 LVS Asn Glv Trp Met Glu Gln Ile Ser Glv Lys Gly Phe Ser Glv 245 250 255 Phe Gln Leu Cys Phe Pro Tyr Tyr Pro Ser Pro Gly Val Leu 260 265 Phe Pro Lys Lys Glu Pro Asp Asp Ser Arg Asp Glu Asp Glu Asp 275 280 285 Glu Asp Glu Ser Ser Glu Glu Asp Ser Glu Asp Glu Glu Pro Pro 290 295 Pro Lys Arg Arg Leu Gln Lys Lys Thr Pro Ala Lys Ser Pro Gly 305 310 315 Lys Ala Ala Ser Val Lys Gln Arg Gly Ser Lys Pro Ala Pro Lys 320 325 330 Val Ser Ala Ala Gln Arg Gly Lys Ala Arg Pro Leu Pro Lys Lys 335 340 345

PCT/US00/02237

Ala Pro Pro Lys Ala Lys Thr Pro Ala Lys Lys Thr Arg Pro Ser 350 355 360 Ser Thr Val Ile Lvs Lvs Pro Ser Glv Glv Ser Ser Lvs Lvs Pro 365 370 375 Thr Ser Ala Arg Lys Glu Val Lys Leu Pro Gly Lys Gly Lys 380 385 390 Ser Thr Met Lys Lys Ser Phe Arg Val Lys Lys

<210> 19 <211> 264 <212> PRT

<213> Homo sapiens

<220> <221> misc-feature

<223> Incyte ID No.: 2572892CD1

<400> 19 Met Pro Arg Ser Phe Leu Val Arg Lys Pro Ser Asp Pro Asn Arg 5 1 10 15 Pro Asn Tyr Ser Glu Leu Gln Asp Ser Asn Pro Glu Phe Thr 20 25 Phe Gln Gln Pro Tvr Asp Gln Ala His Leu Leu Ala Ala Ile Pro 35 40 45 Pro Pro Glu Ile Leu Asn Pro Thr Ala Ser Leu Pro Met Leu Ile 50 55 Trp Asp Ser Val Leu Ala Pro Gln Ala Gln Pro Ile Ala Trp Ala 65 70 75 Leu Arg Leu Gln Glu Ser Pro Arg Val Ser 80 85 90 Leu Ser Asp Glu Asp Ser Glv Lvs Glv Ser Gln Pro Pro Ser Pro 95 100 105 Pro Ser Pro Ala Pro Ser Ser Phe Ser Ser Thr Ser Ala Ser Ser 110 Leu Glu Ala Glu Ala Tyr Ala Ala Phe Pro Gly Leu Gly Gln Val 125 130 135 Lys Gln Leu Ala Gln Leu Ser Glu Ala Lys Asp Leu Gln Ala 140 145 150 Arg Lys Ala Phe Asn Cys Lys Tyr Cys Asn Lys Glu Tyr Leu Ser 155 160 165 Leu Gly Ala Leu Lys Met His Ile Arg Ser His Thr Leu Pro CVS 170 175 180 Cys Gly Thr Cys Gly Lys Ala Phe Ser Arg Pro Trp Leu Leu 185 190 195 Gln Gly His Val Arg Thr His Thr Gly Glu Lys Pro Phe Ser Cvs 200 205 210 Pro His Cys Ser Arg Ala Phe Ala Asp Arg Ser Asn Leu Arg Ala 220 215 225 His Leu Gln Thr His Ser Asp Val Lys Lys Tyr Gln Cys Gln Ala 230 235 240 Cvs Ala Arg Thr Phe Ser Arg Met Ser Leu Leu His Lvs His Gln 245 250 255 Glu Ser Gly Cys Ser Gly Cys Pro Arg 260

<210> 20 <211> 153

<210> 21 <211> 243 PCT/US00/02237

```
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incvte ID No.: 2785674CD1
<400> 20
Met Thr Lys Ile Lys Ala Asp Pro Asp Gly Pro Glu Ala Gln Ala
                                      10
Glu Ala Cys Ser Gly
                    Glu Arg Thr Tyr Gln Glu Leu Leu Val Asn
                 20
                                      25
                                                           30
Gln Asn Pro Ile Ala Gln Pro Leu Ala Ser Arg Arg Leu Thr Arg
                                      40
                                                           45
                 35
                    Ile Lys Lys Ala Val
                                         Lys Gln Lys Gln Ile
Lys Leu Tyr Lys Cys
                 50
                                      55
Arg Arg Gly Val LVs
                    Glu Val Gln Lvs Phe
                                         Val Asn Lys Gly Glu
                                      70
                 65
Lys Gly Ile Met Val Leu Ala Gly Asp Thr Leu Pro Ile Glu
                                                          Val
                                                           90
                 80
                                      85
Tyr Cys His Leu Pro Val Met Cys Glu Asp Arg Asn Leu Pro Tyr
                 95
                                     100
                    Lys Thr Asp Leu Gly Ala Ala Ala Gly Ser
Val Tvr Ile Pro
                Ser
                                     115
                                                          120
                110
                    Val Ile Met Val Lys Pro His Glu Glu Tyr
Lvs Arg Pro Thr
                CVS
                                     130
                                                          135
                125
Gln Glu Ala Tyr Asp Glu Cys Leu Glu Glu Val Gln Ser Leu Pro
                                                          150
                140
                                     145
Leu Pro Leu
```

```
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2797479CD1
```

<400> 21 Met Gly Asp Asp Ile Ser Thr His Ile Ala Pro Lys Glu Leu Arg 15 10 His Lys His Pro Ser Ser Val Asp Glu Val Lys 25 30 20 Gln Leu Arg Ile Ile Leu Gln Gln Gln Val Arg Pro Gly Glu Ser 40 45 35 Thr Val Leu Ala Leu Arg Pro Asn Val Gln Gln Ile Glu Ala Pro 55 50 Asp Val Ser Arg Asp Pro Arg Val Leu Gly Phe Asp Phe Pro Gly 70 75 65 Ser Ala Arg Ile His Glu Gly Thr His Thr Leu Glu Lys Pro Tvr 80 85 90 Phe Glu Cys Lys Gln Cys Gly Lys Leu Leu Ser His Arg Ser Ser 95 100 Arg Arg His Met Met Ala His Thr Gly Asp Gly Pro His Lys Cys 110 115 120 Thr Val Cys Gly Lys Ala Phe Asp Ser Pro Ser Val Phe Gln Arg

WO 00/44900

PCT/US00/02237

				125					130					135
His	Glu	Arg	Thr		Thr	Gly	Glu	Lys	Pro	Tyr	Glu	Cys	Lys	Gln 150
		_		140						_	_	_		
Cys	GIÀ	Lys	Ala		Arg	Thr	ser	ser		Leu	Arg	ьуs	His	
				155					160					165
Thr	Thr	His	Thr	Gly	Glu	Gln	Pro	Tyr	Lys	Cys	Lys	Cys	Gly	
				170					175					180
Ala	Phe	Ser	Asp	Leu	Phe	Ser	Phe	Gln	Ser	His	Glu	Thr	Thr	His
				185					190					195
Ser	Glu	Glu	Glu	Pro	Tyr	Glu	CVS	LVS	Glu	ĊVS	Glv	Lvs	Ala	Phe
				200	~,-		-,-	-1-	205	-,-		-,-		210
Com	Com	Phe	T 110		Dho	0.10	7.~~	uic		7	Thr	Uic	202	Glu
ser	ser	File	ьуь	215	File	Cys	Arg	nis	220	Arg	TILL	HIS	Ser	225
	_	_	_		_									
Glu	Lys	Ser	Tyr		Cys	Gin	Tle	Cys		rys	Leu	ser	vai	
				230					235					240
Ser	Val	Thr												

<400> 22 Met Arg Asp Asn Arg Ala Val Ser Leu Cys Gln Gln Glu Trp Met 10 Cys Pro Gly Pro Ala Gln Arg Ala Leu Tyr Gln 20 30 Arg Lys Asp Ser His Thr Gly Val Gly 35 40 45 Thr Leu Leu Ala Ile Leu Ser Ser Ser Gln Tvr Glu Glu Thr Lvs 50 55 60 Phe Tyr Gly Lys Leu Gln Thr Cys Gln Gln Asn Ser Gln Ile Tyr 75 65 70 Arg Ala Met Ala Glu Gly Leu Trp Glu Gln Gly Phe Leu Arg Thr 80 85 90 Pro Glu Gln Cys Arg Thr Lys Phe Lys Ser Leu Gln Leu Ser Tyr 95 100 105 Arg Lys Val Arg Arg Gly Arg Val Pro Glu Tyr 120 110 115 Glu Glu Met Asn Ala Leu Ser Gly Ser Trp Ala Ser Ala Pro Pro 125 130 Met Ala Ser Asp Ala Val Pro Gly Gln Glu Gly Ser Asp Ile Glu 140 145 Ala Gly Glu Leu Asn His Gln Asn Gly Glu Pro Thr Glu Val Glu 160 165 Asp Gly Thr Val Asp Gly Ala Asp Arg Asp Glu Lys Asp Phe Ara 170 175 Asn Pro Gly Gln Glu Val Arg Lys Leu Asp Phe 190 195 185 Pro Asn Arg Leu Gly Phe Glu Phe Lys Asn Glu Ile Lys Lys Glu 200 205 210 Asn Leu Lys Trp Asp Asp Ser Glu Glu Val Glu Ile Asn Lys Ala 215 220 Leu Gln Arg Lys Ser Arg Gly Val Tyr Trp His Ser Glu Leu Gln

<210> 22

<211> 485 <212> PRT

<213> Homo sapiens

<220>

<221> misc-feature

<223> Incyte ID No.: 2960640CD1

```
235
                 230
                                                            240
Lvs Glv Leu Glu Ser Glu Pro Thr Ser Arg Arg Gln Cys Arg Asn
                                      250
                 245
Ser Pro Gly Glu Ser Glu Glu Lys Thr
                                      Pro
                                          Ser Gln Glu Lys
                                                           Met
                 260
                                      265
                                                           270
Ser His Gln Ser Phe
                    Cys Ala Arg Asp Lys
                                                           Tle
                                      280
                 275
                                                           285
Leu Cys Gly Lys Asn Cys Ser Gln Ser Val His Ser Pro His
                                                           Lvs
                 290
                                      295
Pro Ala Leu Lys Leu Glu Lys Val Ser Gln Cys
                                                           Gly
                305
                                      310
                                                           315
Lys Thr Phe Ser Arg
                    Ser Ser Tyr Leu Val Arg His Gln Arg
                                      325
                320
                                                           330
His Thr Gly Glu Lys Pro His Lys Cys Ser Glu Cys Gly Lys Gly
                                      340
                                                           345
                    Asn Leu Thr Ala His Leu Arg Thr His
                                                           Thr
    Ser Glu Arg Ser
                350
                                      355
                                                           360
Gly Glu Arg Pro Tyr Gln Cys Gly Gln Cys Gly Lys Ser Phe
                                                           Asn
                365
                                      370
   Ser Ser Ser Leu
                    Ile Val His Gln Arg Thr His Thr Gly
                                                           Glu
                380
                                      385
                                                           390
    Pro Tyr Gln Cys
                     Ile Val Cys Gly Lys
                                     400
                395
                                                           405
Ser Gln Phe Ser Ala His Arg Arg Ile His
                                         Thr Gly Glu Ser
                                                           Pro
                410
                                     415
                                                           420
Tyr Lys Cys Ala Val Cys Gly Lys Ile Phe Asn Asn Ser Ser His
                425
                                     430
                                                           435
Phe Ser Ala His Arg Lys Thr His Thr Gly Glu Lys Pro
                                                          Arg
                440
                                     445
                                                           450
Cys Ser His Cys Glu Arg Gly Phe Thr
                                     Lys Asn Ser Ala Leu
                                                          Thr
                                     460
                455
                                                           465
Arg His Gln Thr Val His Met Lys Ala Val
                                         Leu Ser Ser Gln Glu
                                     475
                470
                                                           480
Gly Arg Asp Ala Leu
                485
```

```
<210> 23
<211> 160
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
```

<223> Incyte ID No.: 3454051CD1

<400> 23 Met Ser Trp Thr Cys Pro Arg Cys Gln Gln Pro Val Phe Phe Ala 10 15 Glu Lvs Val Ser Ser Leu Gly Lys Asn Trp His Arg Phe Cys Leu 20 25 30 Lvs Cvs Glu Arg Cvs His Ser Ile Leu Ser Pro Gly Gly His Ala 35 40 Glu His Asn Gly Arg Pro Tyr Cys His Lys Pro Cys Tyr Gly Ala 50 55 60

Leu Phe Gly Pro Arg Gly Pro Pro His Met Lys Thr Phe Thr Gly
65 70 75
Glu Thr Ser Leu Cys Pro Gly Cys Gly Glu Pro Val Tyr Phe Ala
80 85 90

一 稳定键 医硫酸 经净净净 華

PCT/US00/02237

Glu Lys Val Met Ser Leu Gly Arg Asn Trp His Arg Pro Cys Leu 95 100 105 Arg CVS Gln Arg CVs His Lys Thr Leu Thr Ala Gly Ser His Ala 110 115 Pro Tyr Cys His Val Pro Cys Tyr Gly Tyr Glu His Asp Glv Val 135 125 Leu Phe Gly Pro Lys Gly Val Asn Ile Gly Asp Val Gly Cys 140 145 150 Val Lys Ile Lys Phe Lys Ile Tvr Asp Pro 155 160

<210> 24 <211> 511 <212> PRT

<213> Homo sapiens

<220> <221>

<223> Incyte ID No.: 3510640CD1

Met Gln Glu Leu Tyr Ser Thr Pro Ala Ser Arg Leu Asp Ser Phe 10 15 va 1 Val Ala Gln Trp Leu Gln Pro His Arg Glu Trp Lys Glu Glu 20 25 30 Thr Val Glu Glu Phe Leu Arg Gln Glu His Leu Asp Ala Val Ara 45 40 35 Phe Gln Gly Lys Arg Gly Leu Asp Gin Asp Val Arg Val Leu Lvs 5 ō 55 Val Val Lvs Val Glv Ser Phe Gly Asn Gly Thr Val Leu Arg Ser 75 65 70 Ser Cys Phe His Ser Thr Arg Glu Val Glu Leu Val Ala Phe Leu 85 90 80 Val Leu Arg Leu Ile Phe Gln Glu Ala Ala Lvs His His Lvs Asp 95 100 Trp Lys Thr Met Trp Gln Ser Gln Asp Leu Leu Asp Leu Gly Leu 120 110 115 Glu Asp Leu Arg Met Glu Gln Arg Val Pro Asp Ala Leu Val Phe 125 130 135 Thr Ile Gln Thr Arg Gly Thr Ala Glu Pro Ile Thr Val Thr Ile 150 140 145 Val Pro Ala Tyr Arg Ala Leu Gly Pro Ser Leu Pro Asn Ser Gln 165 155 160 Pro Pro Pro Glu Val Tyr Val Ser Leu Ile Lys Ala Cys Gly Gly 170 180 Pro Phe Phe Ser Glu Leu Gln Arg Asn Phe Pro Gly Asn Phe Cys 190 185 Val Lys His Arg Pro Thr Lys Leu Lys Ser Leu Leu Arg Leu Val 200 205 210 Lys His Trp Tyr Gln Gln Tyr Val Lys Ala Arg Ser Pro Arg Ala 215 220 Asn Leu Pro Pro Leu Tyr Ala Leu Glu Leu Leu Thr Ile Tyr Ala 235 230 Trp Glu Met Gly Thr Glu Glu Asp Glu Asn Phe Met Leu Asp Glu 255 245 250 Gly Phe Thr Thr Val Met Asp Leu Leu Leu Glu Tyr Glu Val Ile 260 265 270 Cvs Ile Tyr Trp Thr Lys Tyr Tyr Thr Leu His Asn Ala Ile Ile

PCT/US00/02237

```
275
Glu Asp Cys Val Arg Lys Gln Leu Lys Lys Glu Arg Pro Ile
                                      295
                 290
Leu Asp Pro Ala Asp Pro Thr Leu Asn Val
                                                           315
                                      310
                Ala Gln Arg Ala Ser Gln Cys Leu Lys Gln Asp
    Asp Ile Val
                                      325
                                                           330
                 320
                                                           Val
Cys Cys Tyr Asp Asn Arg Glu Asn Pro Ile
                 335
                                      340
                    Ile His Leu Thr Val
                                                           Tyr
                Asp
Lys Arg Ala Arg
                                                           360
                 350
                                      355
                                         Glu Pro Ile Arg Lys
                    Ile Val Asn Pro Tyr
Pro Asp Phe Asn Leu
                                                           375
                                      370
                 365
                 Ile Arg Arg Thr Arg Gly
Val Lvs Glu Lvs
                                                           390
                                      385
                 380
Arg Leu Ser Phe
                Gln Val Pro Gly Ser Glu
                                                           Ser
                                      400
                                                           405
                 395
                Leu Ala Lys Tyr Gly Ile
                                         Phe Ser His Thr
                                                          His
Ser Arg Cys Ser
                                                           420
                 410
                                      415
                Glu Thr Ile Pro Ser Glu
Ile Tvr Leu Leu
                 425
                                      430
                                                           435
                    Gly Ser Tyr Ala Tyr Ala Ile Asn Pro Asn
Lvs Asn Pro Asp Glv
                 440
                                      445
Ser Phe Ile Leu Gly Leu Lys Gln Gln Ile Glu Asp Gln Gln Gly
                                      460
                 455
Leu Pro Lys Lys Gln Gln Gln Leu Glu Phe Gln Gly Gln Val Leu
                 470
                                      475
Gln Asp Trp Leu Gly Leu Gly Ile Tyr Gly
                                                           Asp
                                      490
                 485
Thr Leu Ile Leu Ser Lys Lys Lys Gly Glu Ala Leu Phe Pro
                                                           510
                 500
                                      505
Ser
```

```
<210> 25
<211> 310
<212> PRT
<213> Homo sapiens
```

Met Arg Pro Leu Gln Ile Val Pro Ser Arg Leu Ile Ser Gln Leu 15 10 1 Tyr Cys Gly Leu Lys Pro Pro Ala Ser Thr Arg Asn Gln Ile Cys 25 30 20 Leu Lys Met Ala Arg Pro Ser Ser Ser Met Lys 35 40 Phe Ala Lys Ala Lys His Ile Val Ile 60 50 55 Ser Ala Glu Ser Gly Val Pro Thr Phe Arg Gly Ala Gly Gly 70 65 Tyr Trp Arg Lys Trp Gln Ala Gln Asp Leu Ala Thr Pro Leu Ala 85 80 Ala His Asn Pro Ser Arg Val Trp Glu Phe Tyr His 105 95 100 Arg Glu Val Met Gly Ser Lys Glu Pro Asn Ala Gly His Arg Ala 115 110

<220> -<223> Incyte ID No.: 3815083CD1

PCT/US00/02237

```
Ile Ala Glu Cys Glu Thr Arg Leu Gly Lys Gln Gly Arg Arg Val
                 125
                                      130
                                                           135
Val Val Ile Thr Gln Asn Ile Asp Glu Leu His Arg Lys Ala Gly
                                      145
                 140
Thr Lys Asn Leu Leu Glu Ile His Gly Ser
                                                      Thr Arg
                                                           165
                 155
                                      160
Cys Thr Ser Cys Gly Val Val Ala Glu Asn Tyr Lys Ser Pro
                 170
                                      175
                                                           180
                     Gly Lys Gly Ala Pro Glu Pro Gly Thr Gln
   Pro Ala Leu Ser
                 185
                                      190
                                                           195
Asp Ala Ser Ile Pro
                    Val Glu Lys Leu Pro Arg Cys Glu Glu Ala
                                                           210
                 200
                                      205
Gly Cys Gly Gly Leu Leu Arg Pro His Val
                                         Val Trp Phe Gly Glu
                                                           225
                 215
                                      220
Asn Leu Asp Pro Ala Ile Leu Glu Glu Val
                                         Asp Arg Glu Leu Ala
                                                           240
                 230
                                      235
His Cys Asp Leu Cys
                    Leu Val Val Glv Thr Ser Ser Val Val
                                                          Tvr
                                      250
                                                           255
                 245
Pro Ala Ala Met Phe Ala Pro Gln Val Ala Ala Arg Gly
                                                          Pro
                 260
                                     265
                                                           270
Val Ala Glu Phe Asn Thr Glu Thr Thr Pro Ala Thr Asn Arg
                                                          Phe
                 275
                                      280
                                                           285
                Gln Gly Pro Cys Gly Thr Thr Leu Pro Glu Ala
Arg Phe His Phe
                 290
                                      295
                                                          300
Leu Ala Cys His Glu Asn Glu Thr Val
                                     Ser
                 305
```

```
<210> 26
```

155

170

1 4:

Ala Glu Glv Lvs Arg Val Cvs Val Ile Asp Ala Ala Val Leu Leu

175 29/91

160

165

180

<211> 331 <212> PRT

<213> Homo sapiens

<220> -<221> misc-feature

<223> Incyte ID No.: 3988457CD1

<400> 26 Met Ala Ile Asn Arg Phe Arg Leu Glu Asn Asp Leu Glu Glu Leu 10 Ala Leu Tyr Gln Ile Gln Leu Leu Lys Asp Leu Arg His Thr Glu 20 25 30 Asn Glu Glu Asp Lvs Val Ser Ser Ser Ser Phe Arg Gln Arg Met 35 40 45 Arg Pro Pro Tyr Glu Arg Pro Glu Leu Pro Leu Gly Asn Leu Leu 50 55 60 Ile Ser Gly Ser Gly Thr Cys Leu Tyr Val Ile Gly Leu Thr Gly 70 65 75 Leu Gly Ala Phe Lvs Ser Ser Ile Ala Gln Arg Leu Lys Gly Val 90 80 85 Ile Asp Ser Asp His Leu Gly His Arg Ala Tyr Ala Pro Gly Gly 95 100 105 Pro Ala Tyr Gln Pro Val Val Glu Ala Phe Gly Thr Asp Ile Len 110 115 His Lvs Asp Glv Ile Ile Asn Arg Lys Val Leu Gly Ser Arg Va1 135 125 130 Lys Gln Leu Lys Ile Leu Thr Asp Ile Met Phe Gly Asn Lys 145 150 140 Pro Ile Ile Ala Lys Leu Ala Arg Glu Glu Met Asp Arg Ala Val

PCT/US00/02237

```
Glu Ala Gly Trp Gln Asn Leu Val His Glu Val Trp Thr Ala Val
                 185
                                      190
                                                           195
Ile Pro Glu Thr Glu Ala Val Arg Arg Ile Val Glu Arg Asp
                                                           Glv
                 200
                                      205
                                                           210
Leu Ser Glu Ala Ala Ala Gln Ser Arg Leu Gln Ser Gln Met
                                                           Sar
                 215
                                      220
Glv Gln Gln Leu Val Glu Gln Ser His Val
                                         Val Leu Ser Ser
                                                           Pro
                 230
                                      235
                                                           240
Cys Gly Ser Arg Ile
                    Ser Pro Asn Ala Arg
                                         Tro Arg Lvs Pro
                                                          Glv
                 245
                                     250
                                                           255
Pro Ser Cys Arg Ser Ala Phe Pro Arg Leu
                                         Ile Arg Pro Ser
                                                          Thr
                 260
                                     265
Glu Lys Phe Ser
                Val Gly Pro Asp Trp Leu Leu Glu Leu Thr
                                                          Ser
                275
                                     280
                                                           285
Asp Pro Val Val
                Arg Arg Asn Gly Gly Leu Asp Ala His Pro
                                                          Glv
                290
                                     295
                                                          300
Ser Gly Pro Glu Val Gln Ala Ile Leu Cys Arg Thr Trp Pro Gly
                305
                                     310
Leu Val Asp Thr Gly Ser Leu Pro Asn Thr Leu Val Phe Gly Gln
                320
                                     325
His
```

Ti

<400> 27 Met Met Thr Ala Glu Ser Arg Glu Ala Thr Gly Leu Ser Pro Gln 10 15 Ala Gln Glu Lys Asp Gly Ile Val Ile Val Lys Val Glu Glu 20 25 30 Glu Asp Glu Glu Asp His Met Trp Gly Gln Asp Ser Thr Leu Gln 35 45 Asp Thr Pro Pro Pro Asp Pro Glu Ile Phe Arg Gln Arg Phe Arg 50 55 60 Arg Phe Cys Tyr Gln Asn Thr Phe Gly Pro Arg Glu Ala Leu Ser 65 70 75 Arg Leu Lys Glu Leu Cys His Gln Trp Leu Arg Pro Glu Ile Asn 80 85 Thr Lys Glu Gln Ile Leu Glu Leu Leu Val Leu Glu Gln Phe Leu 95 100 105 Ser Ile Leu Pro Lys Glu Leu Gln Val Trp Leu Gln Glu Tvr Arg 110 115 120 Pro Asp Ser Gly Glu Glu Ala Val Thr Leu Leu Glu Asp Leu Glu 125 130 135 Leu Asp Leu Ser Gly Gln Gln Val Pro Gly Gln Val His Gly Pro 140 145 Glu Met Leu Ala Arg Gly Met Val Pro Leu Asp Pro Val Gln Glu 155 160 Ser Ser Ser Phe Asp Leu His His Glu Ala Thr Gln Ser His Phe 170 175 180 Lys His Ser Ser Arg Lys Pro Arg Leu Leu Gln Ser Arg Gly Lys 185 190 195 Lys Gln Gly Phe Ile

<210> 27 <211> 200

<212> PRT

<213> Homo sapiens

^{.}

<221> misc-feature

<223> Incyte ID No.: 131890CD1

PCT/US00/02237

200

<210> 28
<211> 100
<212> PRT
<213> Homo sapiens

<220>
<221> misc-feature
<223> Incyte ID No.: 238642CD1

<400> 28 Met Gln Lys Pro Cys Lys Glu Asn Glu Gly Lys Pro Lys Cys Ser 10 1 5 Pro Lys Arg Glu Glu Lys Arg Pro Val 25 20 Gln Gln Thr Glu Gly Asn Phe Arg Gln Arg Leu Leu Gln Ser Leu 40 35 Glu Glu Phe Lys Glu Asp Ile Asp Tyr Arg His Phe Lys Asp Glu 55 60 50 Glu Met Thr Arg Glu Gly Asp Glu Met Glu Arg Cys Leu Glu Glu 70 65 Ile Arg Gly Leu Arg Lys Lys Phe Arg Ala Leu His Ser Asn His 80 85 Arg His Ser Arg Asp Arg Pro Tyr Pro Ile 95

<210> 29
<211> 528
<212> PRT
<213> Homo sapiens

<220>
<221> misc-feature
<223> Incyte ID No.: 669862CD1

<400> 29 Met Ser Ser Pro Tyr Pro Leu Leu Leu Glu Asn Ser Ile Cys Leu 10 15 Phe Phe His Phe Leu Pro Asp Phe Asn Phe Thr Thr Glu Ser Asn 25 30 20 Lys Leu Ser Ser Glu Lys Arg Asn Tyr Glu Val Asn Ala Tyr His 40 35 Gln Glu Thr Trp Lys Arg Asn Lys Thr Phe Asn Leu Met Arg Phe 50 55 60 Ile Phe Arg Thr Asp Pro Gln Tyr Thr Ile Glu Phe Gly Arg Gln 70 75 65 Gln Arg Pro Lys Val Gly Cys Phe Ser Gln Met Ile Phe Lys Lys 85 80 His Lys Ser Leu Pro Leu His Lys Arg Asn Asn Thr Arg Glu Lys 95 100 105 Ser Tyr Glu Cys Lys Glu Tyr Lys Lys Gly Phe Arg Lys Tyr Leu 115 120 110 His Leu Thr Glu His Leu Arg Asp His Thr Gly Val Ile Pro Tyr 135 125 130 Glu Cys Asn Glu Cys Gly Lys Ala Phe Val Val Phe Gln His Phe

PCT/US00/02237

```
150
                                      145
                 140
                    Ile His Thr Asp Leu Lys Pro Tyr Glu Cys
Tle Ard His Ard Lvs
                                      160
                                                           165
                 155
Asn Gly Cys Glu Lys
                     Ala Phe Arg Phe Tyr
                                          Ser Gln Leu Ile Gln
                                      175
                 170
His Gln Ile Ile His
                                     Pro
                                          Tyr Glu Cys Lys Gln
                                      190
                                                           195
                 185
Cys Gly Lys Ala Phe
                                          Leu Thr Glu His
                                                           Gln
                     Arg Arg His Ser His
                 200
                                      205
                                                           210
                                                           Gly
   Ile His Val
                Gly
                             Pro Phe Glu Cys Lys Glu Cys
                 215
                                      220
                                                           225
Glu Thr Phe Arg Leu
                     Tyr Arg His Met Cys
                                                           TIA
                 230
                                      235
                                                           240
His His Gly Val
                     Pro Tyr Lys Cys Lys Glu Cys Gly Lys
                                                           Ala
                Lvs
                                      250
                                                           255
                 245
                     Ser Leu Tyr Gln His Lys Lys Ile His
                                                           Ser
Phe Gly His Arg Ser
                                      265
                                                           270
                 260
Gly Glu Lys Pro
                Tyr
                     Lys Cys Glu Gln Cys Glu Lys Ala Phe
                                                           17= 1
                                      280
                                                           285
                 275
Arg Ser Tyr Leu Leu Val Glu His Gln Arg Ser His Thr Gly
                                                           Glu
                                      295
                                                           300
                 290
    Pro His Glu Cys Met Glu Cys Gly Lys Ala Phe Ser Lys Gly
                 305
                                      310
                                                           315
                                                           Leu
                    His Lys Arg Ile His
                                          Ser Ser Glu Lvs
Ser Ser Leu Leu Lys
                 320
                                      325
                                                           Gln
TVr Asp Cys Lys
                Asp
                    Cys Gly Lys Ala Phe
                                         Cys Arg Gly Ser
                                      340
                 335
Leu Thr Gln His Gln Arg Ile His Thr Gly
                                                           Glu
                                      355
                                                           360
                350
Cys Lys Glu Cys Gly
                    Lys Thr Phe Lys Leu
                                         His Ser Tyr Leu
                                                           Ile
                                                           375
                 365
                                      370
                                                           Lvs
Gln His Gln Ile
                Ile
                    His Thr Asp Leu Lys
                                      385
                                                           390
                 380
Gln Cys Gly Lys
                Ala
                        Ser Arg Val Gly
                                         Asp Leu Lys Thr
                                                           His
                 395
                                      400
Gln Ser Ile His
                Ala Gly Glu Lys Pro
                                     Tyr
                                          Glu Cys Lys Glu
                                                           CVS
                                      415
                                                           420
                 410
                Arg
                    Leu Asn Ser Gln Leu
                                                           Thr
Gly Lys Thr Phe
                 425
                                      430
                                                           435
                    Lys Pro Tyr Val Cys
                                         Lys Glu Cys Lys
                                                           Lvs
Ile His Thr Glv Leu
                 440
                                      445
                                                           450
Ala Phe Arg Ser
                Ile
                    Ser Gly Leu Ser Gln
                                         His Lys Arg Ile
                                                          His
                                     460
                                                           465
                 455
                    Tyr Glu Cys Lys Glu
                                                           Phe
Thr Gly Glu Lys
                Pro
                                         CVs Asp
                                                           480
                470
                                     475
                    Leu Thr Gln His Glu Thr Ile His Thr
                                                          Gly
Asn Arg Ser Asp Arg
                485
                                     490
                                                           495
Val Lys Pro Gln Lys Cys Lys Glu Cys Gly Lys Ala Phe Ser
                500
                                     505
   Tyr Gln Leu Ser Gln His Gln Arg Phe His His Gly Glu Arg
                                     520
                515
Leu Leu Met
```

<210> 30 <211> 350

<212> PRT <213> Homo sapiens

<220>

<221> misc-feature
<223> Incyte ID No.: 1003663CD1

PCT/US00/02237

```
<400> 30
Met His Pro Ala Ala Phe Pro Leu Pro Val Val Val Ala Ala Val
                    Pro Thr Arg Gly Leu Ile Arg Ala Thr Ser
Leu Trp Gly Ala Ala
                                                            30
                 20
Asp His Asn Ala Ser Met Asp Phe Ala Asp Leu Pro Ala
                                                      Leu Phe
                                       40
                                                            45
                 35
Gly Ala Thr Leu Ser Gln Glu Gly Leu Gln Gly Phe Leu Val Glu
                                       55
                                                            60
                 50
                    Ala Cys Ser Pro Ile Ala Pro Pro Pro Pro
Ala His Pro Asp Asn
                                      70
                                                            75
                    Ser Val Phe Ile Ala Leu Leu Arg Arg Phe
Ala Pro Val Asn Glv
                                                            90
                 80
                                       25
                    Leu Lys Val Leu Asn Ala Gln Lys Ala Gly
Asp Cvs Asn Phe Asp
                 95
                                      100
                                                           105
                    Val His Asn Val Asn Ser Asn Glu Leu Leu
Tyr Gly Ala Ala Val
                                                           120
                110
                                     115
Asn Met Val Trp Asn
                    Ser Glu Glu Ile Gln Gln Gln Ile Trp Ile
                                                           135
                                     130
                125
Pro Ser Val Phe Ile Gly Glu Arg Ser Ser Glu Tyr Leu Arg
                                                          Ala
                                                           150
                140
                                     145
                    Lys Gly Ala Arg Val
                                         Leu Leu Val Pro Asp
   Phe Val Tyr Glu
                                     160
                                                           165
                155
                    Gly Tyr Tyr Leu Ile Pro Phe Thr Gly
                                                          Ile
Asn Thr Phe Pro Leu
                170
                                                           180
                    Leu Ala Met Gly Ala Val Met Ile Ala
                                                          Ara
Val Glv Leu Leu Val
                                                           195
                                     190
                185
Cys Ile Gln His Arg Lys Arg Leu Gln Arg Asn Arg Leu Thr Lys
                                     205
                                                           210
                200
Glu Gln Leu Lys Gln
                    Ile Pro Thr His
                                     Asp
                                                           225
                215
                                     220
                                     Asp Glu Tyr Glu Asp Gly
Gln Tyr Asp Val Cys
                    Ala Ile Cys Leu
                230
                                     235
                                                           240
   Lys Leu Arg Val
                    Leu Pro Cys Ala His
                                         Ala Tyr His Ser Arg
                245
                                     250
                                                           255
                Trp
                    Leu Thr Gln Thr Arg
                                         Lys Thr Cys Pro
                                                          Tle
   Val Asp Pro
                                                           270
                                     265
                    His Arg Gly Pro Gly Asp Glu Asp Gln Glu
Cys Lys Gln Pro Val
                275
                                     280
                                                           285
                    Glm Glu Glu Gly Asp Glu Gly Glu Pro
                                                          Ara
   Glu Thr Gln Glv
                290
                                     295
                                                           300
   His Pro Ala Ser
                    Glu Arg Thr Pro
                                     Leu Leu Gly Ser Ser Pro
                                                           315
                305
                                     370
                    Phe Gly Ser Leu Ala
                                         Pro Ala Pro Leu Val
   Leu Pro Thr Ser
                                                          330
                320
                                     325
   Pro Gly Pro Ser Thr Asp Pro Pro Leu Ser Pro Pro Ser
                                                          Ser
                335
                                     340
                                                          345
Pro Val Ile Leu Val
                350
```

<210> 31

<211> 315 <212> PRT

<213> Homo sapiens

<220>

<221> misc-feature

<223> Incyte ID No.: 1432557CD1

```
<400> 31
Met Ala Ala Leu Gly Val Leu Glu Ser Asp Leu Pro Ser Ala Val
                                       10
        Leu Lys Asn Leu Gln Glu Gln Val Met Ala Val Thr Ala
                  20
                                       25
                                                           3 0
Gln Val Lys Ser Leu Thr Gln Lys Val Gln Ala Gly Ala Tyr Pro
                  35
                                       40
                                                           45
Thr Glu Lys Gly Leu Ser Phe Leu Glu Val
                                          Lys Asp Gln Leu Leu
                  50
                                      55
                                                           60
Leu Met Tyr Leu Met Asp Leu Thr His Leu Ile Leu Asp Lys Ala
                  65
                                      70
Ser Gly Gly Ser Leu Gln Gly His Asp Ala Val Leu Arg Leu Val
                  80
                                      85
Glu Ile Arg Thr Val Leu Glu Lys Leu Arg
                                         Pro Leu Asp Gln
                                                         Lys
                                     100
                                                          105
Leu Lys Tyr Gln Ile Asp Lys Leu Ile Lys
                                         Thr Ala Val Thr
                                                          Gly
                 110
                                     115
                                                          120
Ser Leu Ser Glu Asn Asp Pro Leu Arg Phe Lys Pro His Pro
                                                         Ser
                 125
                                                          135
Asn Met Met Ser
                Lys Leu Ser Ser Glu Asp Glu Glu Glu Asp Glu
                 140
                                     145
Ala Glu Asp Asp Gln Ser Glu Ala Ser Gly Lys Lys Ser Val
                                                          Lys
                 155
                                     160
                                                          165
       Ser Lys
                Lys Tyr Val Pro Pro Arg Leu Val Pro Val His
                 170
                                     175
                                                          180
   Asp Glu Thr Glu Ala Glu Arg Glu Lys Lys Arg Leu Glu Arg
                185
                                     190
                                                          195
Ala Lys Arg Arg Ala Leu Ser Ser Ser Val Ile Arg Glu Leu Lys
                 200
                                     205
                                                          210
Glu Gln Tyr Ser Asp Ala Pro Glu Glu Ile Arg Asp Ala Arg
                                                         His
                 215
                                     220
                                                          225
Pro His Val Thr Arg Gln Ser Gln Glu Asp Gln His Arg
                                                     Ile Aso
                230
                                     235
                                                          240
   Glu Glu Ser Met Met Val Arg Leu Ser Val Ser Lys Arg Glu
                                     250
                245
   Gly Arg Arg Lys Arg Ala Asn Val Met Ser Ser Gln Leu His
                260
                                     265
Ser Leu Thr His Phe Ser Asp Ile Ser Ala Leu Thr Gly Gly
                                                         Thr
                275
                                     280
                                                          285
Val His Leu Asp Glu Asp Gln Asn Pro
                                    Ile Lys Lys Arg Lys
                290
                                     295
                                                         300
Ile Pro Gln Lys Gly Arg Lys Lys Gly Phe Arg Arg Arg Arg
                305
                                     310
```

34/91

- region of the area of the second and

<210> 32 <211> 120

<212> PRT <213> Homo sapiens

<220>

<221> misc-feature <223> 1441770CD1

PCT/US00/02237

Leu Arg Tyr Gln Tyr Leu Glu Glu Leu Val Ser Ser Arg Glu Arg 50 60 Ala Ile Cys Ala Leu Arg Glu Glu Leu Glu Met Tvr Lvs Gln Trp 65 70 Ser Glu Ile Lys Ala Cys Met Ala Met Asp Gln Glv Lvs Ile Pro 80 85 9.0 Leu Leu Thr Gly Glu Glu Gln Asn Lys Ser Gln Gln Asn Ser Ser 95 100 105 Arg His Thr Lys Ala Gly Lys Thr Asp Ala Asn Ser Asn Ser Tro 110 115 120

```
<210> 33
<211> 326
<212> PRT
<213> Homo sapiens
```

<220> <221> misc-feature <223> Incyte ID No.: 1456684CD1

<400> 33 Met Gln Glu Glu Pro Leu Pro Gln Gly Asn Asp Pro Glu Pro 5 10 Gly Asp Ser Pro Leu Gly Leu Cys Gln Ser Glu Cvs Met Glu Met 20 25 30 Ser Glu Val Phe Asp Asp Ala Ser Asp Gln Asp Ser LVS 35 40 45 Pro Trp Arg Pro Tyr Tyr Asn Tyr Lys Pro 50 55 60 Lys Lys Met Arg Lys Val Asn Trp Arg Lvs Glu His Glv 65 70 75 Asn Arg Ser Pro Ser His Lys Cys Lys Tyr Pro Ala Glu Leu Asp 80 85 90 Cys Ala Val Gly Lys Ala Pro Gln Asp Lys Pro Phe Glu Glu Glu 100 105 Glu Thr Lys Glu Met Pro Lys Leu Gln Cys Glu Leu Cys Asp Gly 110 115 120 Asp Lys Ala Val Gly Ala Gly Asn Gln Gly 125 130 Leu Thr Ser Arg Pro Tyr Ala Cys Glu Leu Cys Ala Lys Gln Phe 140 145 150 Gln Ser Pro Ser Thr Leu Lys Met His Met Arg Cys Gly 155 160 165 Glu Lys Pro Tyr Gln Cys Lys Thr Cys Gly Arg Cys Val 170 175 180 Gln Gly Asn Leu Gln Lys His Glu Arg Ile His Leu Gly Leu Lys 185 190 195 Glu Phe Val Cys Gln Tyr Cys Asn Lys Ala Phe Thr Leu Asn Glu 200 205 210 Thr Leu Lys Ile His Glu Arg Ile His Thr Gly Glu Lys Tyr 215 220 225 His Cys Gln Phe Cys Phe Gln Arg Phe Leu Tyr Leu Ser Thr Lys 230 235 240 Arg Asn His Glu Gln Arg His Ile Arg Glu His Asn Gly Lys Gly 245 250 255 Tyr Ala Cys Phe Gln Cys Pro Lys Ile Cys Lys Thr Ala Ala Ala 260 265 Leu Gly Met His Gln Lys Lys His Leu Phe Lys Ser Pro Ser Gln 275 280 285

<210> 34 <211> 106

Asn

<210> 35

PCT/US00/02237

of a religion of the second of the second

```
Glu Glu Lys Ile Gly Asp Val Cys His Glu Asn Ser Asn Pro Leu 290 295 300 Glu Asn Gln His Phe Ile Gly Ser Glu Asp Asn Asp Gln Lys Asp 305 Asn Ile Gln Thr Gly Val Glu Asn Val Val Leu 325 325
```

```
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 1602916CD1
<400> 34
Met Phe Pro Trp Met Arg Pro Gln Ala Ala Pro Gly Arg Arg Arg
 7
                  5
                                      10
                                                           15
Glv Arg Gln Thr Tyr Ser Arg Phe Gln Thr Leu Glu Leu Glu Lys
                 20
                                                           30
Glu Phe Leu Phe Asn Pro Tyr Leu Thr Arg Lys Arg Arg Ile Glu
                 35
                                      40
                                                           45
Val Ser His Ala Leu Ala Leu Thr Glu Arg Gln Val Lys Ile Trp
                 50
                                      55
Phe Gln Asn Arg Arg Met Lys Trp Lys Lys Glu Asn Asn Lys Asp
                 65
                                      70
                                                           75
Lys Phe Pro Val Ser Arg Gln Glu Val Lys Asp Gly Glu Thr Lys
                 80
                                      85
                                                           90
Lys Glu Ala Gln Glu Leu Glu Glu Asp Arg Ala Glu Arg Leu Thr
                 95
                                    100
```

```
<211> 209
<212> PRT '
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 1692816CD1
```

<400> 35 Met Asn Pro Ser Met Lys Gln Lys Gln Glu Glu Ile Lys Glu Asn 1 5 10 15 Ile Lys Asn Ser Ser Val Pro Arg Thr Leu Lys Met Ile Gln 20 Pro Ser Ala Ser Gly Ser Leu Val Gly Arg Glu Asn Glu Leu Ser 35 40 Ala Gly Leu Ser Lys Arg Lys His Arg Asn Asp His Leu Thr Ser 50 55 Thr Thr Ser Ser Pro Gly Val Ile Val Pro Glu Ser Ser Glu Asn 70 65 Lys Asn Leu Gly Gly Val Thr Gln Glu Ser Phe Asp Leu Met Ile 80 85 90 Lys Glu Asn Pro Ser Ser Gln Tyr Trp Lys Glu Val Ala Glu Lys

WO 60/44966

PCT/US00/02237

				95					100					105
Arg i	Arg	Lys	Ala	Leu 110	Tyr	Glu	Ala	Leu	Lys 115	Glu	Asn	Glu	Lys	Leu 120
His l	Lys	Glu	Ile	Glu 125	Gln	Lys	Asp	Asn	Glu 130	Ile	Ala	Arg	Leu	Lys 135
Lys (Glu	Asn	Lys	Glu 140	Leu	Ala	Glu	Val	Ala 145	Glu	His	Val	Gln	Tyr 150
Met 1	Ala	Glu	Leu	11e	Glu	Arg	Leu	Asn	Gly 160	Glu	Pro	Leu	Asp	Asn 165
Phe (Glu	Ser	Leu	Asp 170	Asn	Gln	Glu	Phe	Asp 175	Ser	Glu	Glu	Glu	Thr 180
Val (Glu	Asp	Ser	Leu 185	Val	Glu	Asp	Ser	Glu 190	Ile	Gly	Thr	Cys	Ala 195
Glu	Gly	Thr	Val	Ser 200	Ser	Ser	Thr	Asp	Ala 205	Lys	Pro	Cys	Ile	

<210> 36 <211> 212 <212> PRT

<400> 36 Met Leu Gly Asn Glu Trp Ser Lys Leu Pro Pro Glu Glu Lys Gln 15 Arg Tyr Leu Asp Glu Ala Asp Arg Asp Lys Lys Glu Arg Tyr Met 3 0 20 Glu Leu Glu Gln Tyr Gln Lys Thr Glu Ala Tyr Lys Val Phe Ser 40 35 Arg Gln Lys Gly Lys Ser His Arg Gln Asp Thr Gln Asp 60 50 55 Ala Ala Arg Gln Ala Thr His Asp His Glu Lys Glu Thr Glu Val 75 70 65 Lys Glu Arg Ser Val Phe Asp Ile Pro Ile Phe Thr Glu Glu Phe 85 90 80 Leu Asn His Ser Lys Ala Arg Glu Ala Glu Leu Arg Gln Leu Arg 100 95 Lys Ser Asn Met Glu Phe Glu Glu Arg Asn Ala Ala Leu Gln Lys 120 115 110 His Val Glu Ser Met Arg Thr Ala Val Glu Lys Leu Glu Val Asp 130 135 125 Val Ile Gln Glu Arg Ser Arg Asn Thr Val Leu Gln Gln His Leu 145 Glu Thr Leu Arg Gln Val Leu Thr Ser Ser Phe Ala Ser Met Pro 165 155 160 Leu Pro Gly Ser Gly Glu Thr Pro Thr Val Asp Thr Ile Asp 170 175 180 Tyr Met Asn Arg Leu His Ser Ile Ile Leu Ala Asn Pro Gln Asp 190 195 185 Asn Glu Asn Phe Ile Ala Thr Val Arg Glu Val Val Asn Arg Leu 210 205 200 Asp Arg

<213> Homo sapiens

^{-220&}gt;

<221> misc-feature
<223> Incyte ID No.: 1968191CD1

<210> 37

PCT/US00/02237

```
<211> 359
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2052061CD1
<400> 37
Met Val Asp Met Asp Lys Leu Ile Asn Asn Leu Glu Val Gln Leu
                                       10
                    Ser Met Gln Val Phe Lys Gln Val Thr Ala
Asn Ser Glu Gly Gly
                 20
                                       25
                                                            3 0
    Val Arg Asn Arg Asp Pro Pro Glu Ile Glu Tyr Thr Ser
                                                          Asn
                 35
                                      40
                                                            45
Met Thr Ser Pro Thr Leu Leu Asp Ala Asn Pro Met Glu Asn Pro
                 50
                                      55
                                                            60
Ala Leu Phe Asn Asp Ile Lys Ile Glu Pro Pro Glu Glu Leu Leu
                 65
                                      70
Ala Ser Asp Phe Ser Leu Pro Gln Val Glu Pro Val Asp Leu
                                                          Ser
                 80
                                      85
                                                            90
Phe His Lys Pro Lys Ala Pro Leu Gln Pro Ala Ser Met Leu Gln
                                     100
                 95
                                                          105
   Pro Ile Arg Pro Pro Lys Pro Gln Ser Ser Pro Gln Thr Leu
                                                          120
                110
                                     115
Val Val Ser Thr Ser Thr Ser Asp Met Ser
                                         Thr Ser Ala Asn
                                                          Ile
                125
                                     130
                                                          135
Pro Thr Val Leu Thr Pro Gly Ser Val Leu
                                         Thr Ser Ser Gln
                140
                                     145
                                                          150
Thr Gly Ser Gln Gln Ile Leu His Val Ile
                                         His Thr Ile Pro
                                                          Ser
                155
                                     160
                                                          165
Val Ser Leu Pro Asn Lys Met Gly Gly Leu Lys Thr Ile Pro
                                                          Val
                170
                                     175
                                                          180
Val Val Gln Ser Leu Pro Met Val Tyr Thr
                                         Thr Leu Pro Ala
                                                          Asp
                185
                                     190
Gly Gly Pro Ala Ala Ile Thr Val Pro Leu
                                         Ile Gly Gly Asp
                                                          Gly
                200
                                     205
                                                          210
Lys Asn Ala Gly Ser Val Lys Val Asp Pro Thr Ser Met Ser
                                                          Pro
                215
                                     220
                                                          225
Leu Glu Ile Pro Ser Asp Ser Glu Glu Ser
                                         Thr Ile Glu Ser
                                                          240
    Ser Ala Leu Gln Ser Leu Gln Gly Leu Gln Glu Pro
                                                          Ala
                245
                                     250
                                                          255
   Met Ala Gln Met Gln Gly Glu Glu Ser Leu Asp Leu Lys
                                                          Ara
                260
                                     265
                                                          270
Arg Arg Ile His Gln Cys Asp Phe Ala Gly Cys Ser Lys Val
                                                          Tyr
                275
                                     280
                                                          285
Thr Lys Ser Ser His Leu Lys Ala His Arg Arg Ile His
                                                          Gly
                290
                                     295
                                                          300
Glu Lys Pro Tyr Lys Cys Thr Trp Asp Gly
                                         Cys Ser Trp Lys
                305
                                     310
                                                          315
Ala Arg Ser Asp Glu Leu Thr Arg His Phe Arg Lys His Thr
                                                          Glv
                320
                                     325
                                                          330
Ile Lys Pro Phe Arg Cys Thr Asp Cys Asm Arg Ser Phe Ser Arg
                                     340
                335
Ser Asp His Leu Ser Leu His Arg Arg Arg His Asp Thr Met
                350
                                     355
```

<210> 38

<400> 38

PCT/US00/02237

<211> 445 <212> PRT <213> Homo sapiens

<221> misc-feature <223> Incyte ID No.: 2056207CD1

Met Val Glu Cys Ile Arg Glu Val Asn Glu Val Ile Gln Asn Pro 15 5 10 Arg Ile Leu Leu Ser His Phe Asn Trp Asp Lys Ala Thr Ile Thr 20 30 Glu Arg Tyr Phe Asp Gly Asn Leu Glu Lys Leu 40 45 35 Phe Ala Glu Cys His Val Ile Asn Pro Ser Lys Lys Ser Arg Thr 55 60 50 Arg Gln Met Asn Thr Arg Ser Ser Ala Gln Gln Asp Met Pro Cvs 70 75 65 Tyr Leu Asn Tyr Pro Asn Ser Tyr Phe Thr Gly Leu Glu 80 85 90 Thr Cys Gly His Lys Phe Cys Met Gln Cys Trp 100 105 95 Ile Met Glu Glu Gly Met Gly Gln Thr Ile Ser Cys 120 110 115 Ile Leu Val Asp Asp Asn Thr Val Met Ara Ala His Gly Cys Asp 125 130 135 Ile Thr Asp Ser Lys Val Lys Leu Lys Tyr Gln His Leu 140 145 150 Pro Asn Ser Phe Val Glu Cys Asn Arg Leu Leu Lys 155 160 165 Val Gln Tyr Ala Ala Pro Asp Cys His His Val Val Lys 175 170 180 Pro Val Arg Cys Lys Cys Gly Arg Gln Phe Cys Phe Asn Cys 195 185 190 His Lys CVS Gly Glu Asn Trp Asp Pro Val Lys Lys Trp Leu Lys 210 200 205 Cys Glu Thr Ser Asn Trp Tle Ile Lys Lys Asp Asp Asp 215 220 225 Ala Asn Thr Lys Glu Cys Pro CVS Glu 235 240 230 Val Cys CVS Asp Gly Gly CVS Asn His Met Arg Asn Gln Asn 245 250 255 His Ala Glu Phe Cys Val Cys Leu Gly Pro Trp Glu Pro 260 265 270 Ser Ala Trp Tyr Asn Cys Asn Arg Tyr A1a 280 285 275 Asp Ser Lys Ala Ala Arg Ala Gln Glu Arg Arg Ala Ala Leu 290 295 300 Tyr Cys Asn Arg Tyr Met Asn His Met Gln Ser Tyr Leu Phe 310 315 305 Arg Phe Glu His Lys Leu Tyr Ala Gln Val Lys Gln Lys Met 325 330 320 Glu Glu Met Gln Gln His Asn Met Ser Trp Ile Glu Val Gln Phe 335 340 345 Leu Lys Lys Ala Val Asp Val Leu Cys Gln Cys Arg Ala Thr 360 350 Met Tyr Thr Tyr Val Phe Ala Phe Tyr Leu Lys Asn Asn Gln 370 375 365 Ser Ile Ile Phe Glu Asn Asn Gln Ala Asp Leu Glu Asn Ala Thr 380 385 390

PCT/US00/02237

```
Glu Val Leu Ser Gly Tyr Leu Glu Arg Asp Ile Ser Gln Asp Ser
                                                           405
                395
                                      400
                                                           Cys
   Gln Asp Ile Lys Gln Lys Val Gln Asp
                                         Lys Tyr Arg Tyr
                                                           420
                                      415
                410
                    Val Leu Leu Gln His
                                          Val His Glu Glv
                                                           Tvr
Glu Ser Arg Arg Arg
                                     430
                                                           435
                425
Glu Lys Asp Leu Trp Glu Tyr Ile Glu Asp
                                      445
```

```
<210> 39
<211> 433
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incvte ID No.: 2101803CD1
<400> 39
Met Arg Ala Glu Gly Leu Gly Gly Leu Glu Arg Phe Cys Ser Pro
                                      10
                                                            15
                    Leu Arg Ala Leu Gln
                                         Pro Phe Gln Val Glv
Gly Lys Gly Arg Gly
                 20
                                       25
                                                            30
Asp Leu Leu Phe Ser
                    Cys Pro Ala Tyr Ala
                                                           Val
                                      40
                                                            45
                 35
Asn Glu Arg Gly Asn His Cys Glu Tyr Cys
                                       55
                                                            60
                 50
                                 Lys Gln
                                                           Asn
                                          Ala Phe
    Leu Ser Lys Cys Gly Arg Cys
                 65
                                      70
                                                            75
                                                           Cys
Val Glu Cvs Gln Lvs
                    Glu Asp Trp Pro Met
                                          His Lys
                                                            90
                 80
                                      85
Ser Pro Met Val Val
                    Phe Gly Glu Asn Trp
                                         Asn Pro
                                                  Ser Glu Thr
                                                           105
                                      100
                 95
Val Arg Leu Thr Ala Arg Ile Leu Ala Lvs
                                         Gln Lvs Ile His
                                                           Pro
                                      115
                                                           120
                110
Glu Arg Thr Pro Ser Glu Lys Leu Leu Ala
                                         Val Lys Glu Phe Glu
                                      130
                                                           135
                125
                                                           Gln
   His Leu Asp Lys
                    Leu Asp Asn Glu Lys
                                         Lys Asp Leu Ile
                                     145
                140
Ser Asp Ile Ala Ala Leu His His Phe Tyr
                                                           Glu
                155
                                     160
                                                           165
                    Ser Leu Val Val Leu
                                         Phe Ala Gln Val
                                                           Asn
   Pro Asp Asn Asp
                17°0
                                     175
                                                           180
                    Ile Glu Asp Glu Glu
                                         Leu Ser His Leu Glv
   Asn Gly Phe Thr
                                     190
                                                           195
                185
                Pro Asp Val Ala Leu Met Asn His Ser Cys
                                                           Ċvs
Ser Ala Ile Phe
                                     205
                                                           210
                200
Pro Asn Val Ile Val Thr Tyr Lys Gly Thr Leu Ala Glu Val
                                     220
                                                           225
                215
Ala Val Gln Glu Ile Lys Pro Gly Glu Glu Val Phe Thr Ser
                                                           Tyr
                230
                                     235
                                                           240
Ile Asp Leu Leu Tyr Pro Thr Glu Asp Arg Asn Asp Arg Leu
                                                          Ara
                                     250
                                                           255
                245
                Phe Thr Cys Glu Cys Gln Glu Cys
                                                           Lys
Asp Ser Tyr Phe
                                     265
                260
                    Lys Val Glu Ile Arg Lys Leu Ser Asp
                                                          Pro
Asp Lys Asp Lys Ala
                275
                                     280
                                                           285
```

Pro Lys Ala Glu Ala Ile Arg Asp Met Val Arg Tyr Ala Arg

290

295 40/91 Asn

PCT/US00/02237

```
Val Ile Glu Glu Phe Arg Arg Ala Lys His Tyr Lys Ser Pro Ser
                                     310
Glu Leu Leu Glu Ile Cys Glu Leu Ser Gln Glu Lys Met Ser Ser
                                     325
                320
   Phe Glu Asp Ser Asn Val Tyr Met Leu His Met Met Tyr Gln
                335
                                     340
                                                          3/15
Ala Met Gly Val Cys
                    Leu Tyr Met Gln Asp Trp Glu Gly Ala Leu
                350
                                     355
                                                          360
Gln Tyr Gly Gln Lys
                    Ile Ile Lys Pro Tyr Ser Lys His Tyr Pro
                365
                                     370
   Tyr Ser Leu Asn Val Ala Ser Met Trp
                                         Leu Lys Leu Gly Arg
                380
                                     385
Leu Tyr Met Gly Leu Glu His Lys Ala Ala Gly Glu Lys Ala Leu
                395
                                     400
                                                          405
Lys Lys Ala Ile Ala Ile Met Glu Val Ala His Gly Lys Asp
                                                         His
                410
                                     415
Pro Tyr Ile Ser Glu Ile Lys Gln Glu Ile Glu Ser His
                                     430
```

```
<210> 40
<211> 355
<212> PRT
```

<220> <221> misc-feature

<400> 40 Met Ser Val Asn Tyr Ala Ala Gly Leu Ser Pro Tyr Ala Asp Lys 10 Gly Lys Cys Gly Leu Pro Glu Ile Phe Asp Pro Pro Glu Glu Leu 20 25 30 Glu Arg Lys Val Trp Glu Leu Ala Arg Leu Val Trp Gln Ser Ser 35 40 45 Asn Val Val Phe His Thr Gly Ala Gly Ile Ser Thr Ala Ser Glv 50 55 60 Ile Pro Asp Phe Arg Gly Pro His Gly Val Trp Thr Met Glu Glu 65 70 Arg Gly Leu Ala Pro Lys Phe Asp Thr Thr Phe Glu Ser Ala Arg 80 85 90 Pro Thr Gln Thr His Met Ala Leu Val Gln Leu Glu Arg Val Glv 95 100 105 Leu Leu Arg Phe Leu Val Ser Gln Asn Val Asp Gly Leu His Val 110 115 120 Arg Ser Gly Phe Pro Arg Asp Lys Leu Ala Glu Leu His Gly Asn 125 130 Met Phe Val Glu Glu Cys Ala Lys Cys Lys Thr Gln Tyr Val Arg 140 145 150 Asp Thr Val Val Gly Thr Met Gly Leu Lys Ala Thr Gly Arg Leu 155 160 165 Cys Thr Val Ala Lys Ala Arg Gly Leu Arg Ala Cys Arg Gly Glu 170 175 180 Leu Arg Asp Thr Ile Leu Asp Trp Glu Asp Ser Leu Pro Asp Arg 185 190 195 Asp Leu Ala Leu Ala Asp Glu Ala Ser Arg Asn Ala Asp Leu Ser 200 205 210 Ile Thr Leu Gly Thr Ser Leu Gln Ile Arg Pro Ser Gly Asn Leu 215 220

<213> Homo sapiens

<223> Incyte ID No.: 2112362CD1

PCT/US00/02237

Pro Leu Ala Thr Lys Arg Arg Gly Gly Arg Leu Val Ile Val Asn 230 235 Gln Pro Thr Lys His Asp Arg His Ala Asp Leu Arg Ile His 245 250 255 Tyr Val Asp Glu Val Met Thr Arg Leu Met Lys His Leu Glv 260 265 270 Leu Glu Ile Pro Ala Trp Asp Gly Pro Arg Val Leu Glu Arg Δla 275 280 285 Leu Pro Pro Leu Pro Arg Pro Pro Thr Pro Lys Leu Glu Pro 290 295 300 Glu Glu Ser Pro Thr Arg Ile Asn Gly Ser Ile Pro Ala Gly Pro 305 310 315 Lys Gln Glu Pro Cys Ala Gln His Asn Gly Ser Glu Pro Ala Ser 320 325 Pro Lvs Arg Glu Arg Pro Thr Ser Pro Ala Pro His Arg Pro Pro 335 340 345 Lys Arg Val Lys Ala Lys Ala Val Pro Ser 350 355

```
<210> 41
<211> 443
```

<400> 41

Met Asp Arg Leu Gly Ser Phe Ser Asn Asp Pro Ser Asp Lys Pro 10 15 Pro Cys Arg Gly Cys Ser Ser Tyr Leu Met Glu Pro Tyr Ile Lys 20 25 3.0 Cys Ala Glu Cys Gly Pro Pro Pro Phe Phe Leu Cys Leu Gln Cys 35 40 Phe Thr Arg Gly Phe Glu Tyr Lys Lys His Gln Ser Asp His Thr 50 55 60 Tyr Glu Ile Met Thr Ser Asp Phe Pro Val Leu Asp Pro 65 70 75 Thr Ala Gln Glu Met Ala Leu Leu Glu Ala Val Met Asp Cys 80 85 90 Gly Phe Gly Asn Trp Gln Asp Val Ala Asn Gln Met Cys Thr Lys 95 100 105 Thr Lys Glu Glu Cys Glu Lys His Tyr Met Lys His Phe Ile Asn 110 115 120 Asn Pro Leu Phe Ala Ser Thr Leu Leu Asn Leu Lys Gln Ala Glu 125 130 135 Glu Ala Lys Thr Ala Asp Thr Ala Ile Pro Phe His Ser Thr Asp 140 145 150 Asp Pro Pro Arg Pro Thr Phe Asp Ser Leu Leu Ser Arg Asp Met 155 160 165 Ala Gly Tyr Met Pro Ala Arg Ala Asp Phe Ile Glu Glu Phe Asp 170 175 180 Asn Tyr Ala Glu Trp Asp Leu Arg Asp Ile Asp Phe Val Glu Asp 185 190 195 Asp Ser Asp Ile Leu His Ala Leu Lys Met Ala Val Val Ile 200 205 210 Tyr His Ser Arg Leu Lys Glu Arg Gln Arg Arg Lys Lys Ile Ile 215 220 225

<212> PRT

<213> Homo sapiens

<220>

<221> misc-feature <223> Incyte ID No.: 2117346CD1

PCT/US00/02237

```
Arg Asp His Gly Leu Ile Asn Leu Arg Lys Phe Gln Leu Met Glu
                 230
                                      235
                                                           240
Arg Arg Tyr Pro Lys Glu Val Gln Asp Leu Tyr Glu Thr Met Arg
                 245
                                      250
                                                           255
Arg Phe Ala Arg Ile Val Glv Pro Val Glu His Asp Lvs Phe
                                                          T10
                 260
                                      265
                                                           270
Glu Ser His Ala Leu Glu Phe Glu Leu Arg Arg Glu Ile Lys
                                                          Arg
                                      280
                 275
                                                           285
Leu Gln Glu Tyr Arg
                    Thr Ala Gly Ile Thr Asn Phe Cys Ser
                                                          Ala
                 290
                                      295
                                                           300
Arg Thr Tyr Asp His Leu Lys Lys Thr Arg Glu Glu Glu Arg
                                                           Leu
                 305
                                      310
                                                           315
Lys Arg Thr Met Leu Ser Glu Val Leu Gln Tyr Ile Gln Asp
                 320
                                      325
                                                           330
Ser Ala Cys Gln Gln Trp Leu Arg Arg Gln Ala Asp Ile Asp
                                                          Ser
                 335
                                      340
                                                           345
Glv Leu Ser Pro Ser Ile Pro Met Ala Ser Asn Ser Gly Arg
                                                          Ara
                 350
                                      355
                                                           360
Ser Ala Pro Pro
                Leu Asn Leu Thr Gly Leu Pro Gly Thr Glu
                                                           Lvs
                 365
                                      370
                                                           375
Leu Asn Glu Lys Glu
                    Lys Glu Leu Cys Gln Met Val Arg Leu
                                                          Va1
                 380
                                      385
                                                           390
Pro Glv Ala Tvr
                Leu Glu Tvr Lvs Ser Ala Leu Leu Asn Glu Cvs
                 395
                                      400
                                                           405
Asn Lys Gln Gly Gly Leu Arg Leu Ala Gln Ala Arg Ala Leu
                                                          T1 🗅
                 410
                                      415
Lys Ile Asp Val Asn Lys Thr Arg Lys Ile Tyr Asp Phe Leu
                                                          Ile
                 425
                                     430
                                                           435
Arg Glu Gly Tyr
                Ile Thr Lys Gly
                440
```

```
<210> 42
<211> 164
<212> PRT
```

<220>

<400> 42 Met Thr Ala Ser Ala Gln Pro Arg Gly Arg Arg Pro Gly Val Gly 1 5 10 15 Val Gly Val Val Val Thr Ser Cys Lys His Pro Arg Cys Val Len 20 25 Leu Gly Lys Arg Lys Gly Ser Val Gly Ala Gly Ser Phe Gln Leu 35 40 45 Pro Gly Gly His Leu Glu Phe Gly Glu Thr Trp Glu Glu Cys Ala 50 60 Gln Arg Glu Thr Trp Glu Glu Ala Ala Leu His Leu Lys Asn Val 65 70 75 His Phe Ala Ser Val Val Asn Ser Phe Ile Glu Lys Glu Asn Tyr 80 85 90 His Tyr Val Thr Ile Leu Met Lys Gly Glu Val Asp Val Thr His 95 100 105 Asp Ser Glu Pro Lys Asn Val Glu Pro Glu Lys Asn Glu Ser Trp 110 115 120 Glu Trp 'al Pro Trp Glu Glu Leu Pro Pro Leu Asp Gln Leu Phe 125 130 135

<213> Homo sapiens

<221> misc-feature <223> Incyte ID No.: 2119917CD1

PCT/US00/02237

Trp Gly Leu Arg Cys Leu Lys Glu Gln Gly Tyr Asp Pro Phe Lys 140 145 150 150 Glu Asp Leu Asn His Leu Val Gly Tyr Lys Gly Asn His Leu 155 160

```
<210> 43
<211> 215
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2123456CD1
<400> 43
Met Leu Gly Ala Glu Trp Ser Lys Leu Gln Pro Thr Glu Lys Gln
                                      10
                                                           15
 1
Arg Tyr Leu Asp Glu Ala Glu Arg Glu Lys Gln Gln Tyr Met
                 2.0
                                      25
Glu Leu Arg Ala Tyr Gln Gln Ser Glu Ala Tyr Lys Met Cys
                                                          Thr
                                      40
Glu Lys Ile Gln Glu Lys Lys Ile Lys Lys Glu Asp Ser Ser
                                                           60
                 50
                                      55
Gly Leu Met Asn Thr Leu Leu Asn Gly His Lys Gly Gly Asp Cys
                                      70
                 65
Asp Gly Phe Ser Thr Phe Asp Val Pro Ile Phe Thr Glu Glu Phe
                                      85
                                                           90
                 20
Leu Asp Gln Asn Lvs Ala Arg Glu Ala Glu Leu Arg Arg Leu Arg
                                     100
                                                          105
                 95
Lys Met Asn Val Ala Phe Glu Glu Gln Asn Ala Val Leu Gln
                                                         Arg
                110
                                     115
                                                          120
His Thr Gln Ser Met Ser Ser Ala Arg Glu Arg Leu Glu Gln Glu
                125
                                     130
                                                          135
Leu Ala Leu Glu Glu Arg Arg Thr Leu Ala Leu Gln Gln Gln Leu
                                                          150
                                     145
                140
Gln Ala Val Arg Gln Ala Leu Thr Ala Ser Phe Ala Ser Leu
                                                         Pro
                                     160
                                                          165
                155
Val Pro Gly Thr Gly Glu Thr Pro Thr Leu Gly Thr Leu Asp Phe
                                     175
                                                          180
                170
Tyr Met Ala Arg Leu His Gly Ala Ile Glu Arg Asp Pro Ala Gln
                                     190
                                                          195
                185
His Glu Lys Leu Ile Val Arg Ile Lys Glu Ile Leu Ala Gln Val
                200
                                     205
                                                          210
```

```
<210> 44
<211> 539
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2148792CD1
```

215

Ala Ser Glu His Leu

<400> 44

PCT/US00/02237

Met Ala Ala Leu Phe Leu Ser Ala Pro Pro Gln Ala Glu Val Thr 10 Ala Val Tyr Leu Ser Arg Glu Glu Trp Gly Arg Phe Glu Asp Val 30 25 Asp Val Met Leu Glu Gly Pro Ala Gln Arg Gly Leu Tvr Arg 40 35 Thr Tyr Gly Asn Leu Val Ser Leu Gly Val Gly Pro Ala Gly Pro 55 50 Gly Asp Glu Pro Trp Pro Gly Val Ile Ser Gln Leu Glu Arg 75 70 65 Glu His Leu Arg Val Leu Asp Val Gln Gly Thr Ser Gly Lys 90 80 Asn Ser Pro Ala Leu Gly Thr Arg Thr Glu Tyr Lys Glu Leu Thr 100 95 Gln Glu Thr Phe Gly Glu Glu Asp Pro Gln Gly Ser Glu Pro 120 110 115 His Ile Ser Lys Ser Glu Gly Ser Leu Glu Glu Ala Cys Asp 125 130 Leu Val Glu Gln Arg Gly Pro Arg Ala Val Thr Leu Thr Agn 145 150 140 Glu Ser Ser Arg Glu Ser Gly Gly Asn Leu Arg Leu Leu Ser 165 155 160 Pro Val Pro Asp Gln Arg Pro His Lys Glu Cys Asp Ile 175 180 170 Val Gln Ser Phe Glu Gln Arg Ser Tyr Leu Asn 190 195 185 Asn Ser Gly Glu Ile Thr Asn Thr Val Arg His Ard Ser Lvs Lvs 210 200 205 Thr Ser Ala Asn Leu Val Val Lys Glu Asp Gln Lys Ile Pro 220 225 215 Tyr Cys Ser Tyr Cys Gly Lys Thr Phe Arg Lys Lys Leu His 240 235 230 Ser Ala Asn Leu Val Lys His Gln Arg Leu His Thr Glu Glu 250 245 Asp Glu Cys Gly Lys Ala Phe Ser Gln Ser Lys Pro Tyr Lys Cys 265 270 260 Glu Phe Ile Asn His Arg Arg Met His Ser Gly Glu Ile Pro 275 280 Arg Cys Asp Glu Cys Gly Lys Thr Phe Thr Arg Arg Pro Asn 300 290 295 Met Lys His Gln Arg Ile His Thr Gly Lys 315 305 310 Lys His Phe Ser Ala Tyr Ser Ser Leu Tle Cys Gly Glu Cys Gly 325 320 Tyr His Gln Arg Ile His Thr Gly Glu Lys Pro Tyr Lys Asn 345 340 335 Phe Ser Asp Gly Ser Ile Leu Ile Arg His Asp Cys Gly Lys Ala 360 350 355 Arg Arg Thr His Thr Gly Glu Lys Pro Phe Glu Cys Lys Glu Cys 375 370 365 Gly Lys Gly Phe Thr Gln Ser Ser Asn Leu Ile Gln His Gln Arg 390 380 385 Thr Gly Glu Lys Pro Tyr Lys Cys Asn Glu Cys Glu Lys 405 395 400 Thr Lys Leu Val Glu His Gln Arg Ser His Ile Gln Lvs 420 415 410 Thr Gly Glu Lys Pro Tyr Glu Cys Asn Asp Cys Gly Lys Val Phe 435 430 425 Leu Ile Gln His Gln Arg Ile His Thr Gly Ser Gln Ser Thr His 450 440 445 Glu Lys Pro Tyr Lys Cys Ser Glu Cys Gly Lys Ala Phe His Asn 465 460 455

480

Thr

135

Glu 150

WO 00/44900

470

PCT/US00/02237

```
Pro Tyr Arg Cys Ser Asp Cys Lys Lys Ala Phe Ser Gln Ser Thr
                485
                                     490
Tyr Leu Ile Gln His Arg Arg Ile Hıs Thr Gly Glu Lys Pro Tyr
                                                          510
                500
                                     505
Lys Cys Ser Glu Cys Gly Lys Ala Phe Arg His Ser Ser Asn Met
                                     520
                                                          525
                515
Cys Gln His Gln Arg Ile His Leu Arg Glu Asp Phe Ser Met
                                     535
                530
<210> 45
<211> 182
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2751943CD1
<400> 45
Met Ala Arg Leu Leu Trp Leu Leu Arg Gly Leu Thr Leu Gly Thr
                                                           15
                                      10
  1
                  5
Ala Pro Arg Arg Ala Val Arg Gly Gln Ala Gly Gly Gly Pro
                                      25
                                                           30
                 20
Gly Thr Gly Pro Gly Leu Gly Glu Ala Gly Ser Leu Ala Thr Cys
                 35
                                      40
                                                           45
Glu Leu Pro Leu Ala Lys Ser Glu Trp Gln Lys Lys Leu Thr Pro
                                      55
                                                           60
                 50
Glu Gln Phe Tyr Val Thr Arg Glu Lys Gly Thr Glu Pro Pro Phe
                                      70
                                                           75
                 65
Ser Gly Ile Tyr Leu Asn Asn Lys Glu Ala Gly Met Tyr His Cys
                                                           90
                 80
                                      85
                                                         Cys
Val Cys Cys Asp Ser Pro Leu Phe Ser Ser Glu Lys Lys Tyr
                                                         105
                 95
                                     100
Ser Gly Thr Gly Trp Pro Ser Phe Ser Glu Ala His Gly Thr Ser
                                     115
                                                          120
                110
```

Gly Ser Asp Glu Ser His Thr Gly Ile Leu Arg Arg Leu Asp

Ser Leu Gly Ser Ala Arg Thr Glu Val Val Cys Lys Gln Cys

Ala His Leu Gly His Val Phe Pro Asp Gly Pro Gly Pro Asn Gly

Gln Arg Phe Cys Ile Asn Ser Val Ala Leu Lys Phe Lys Pro Arg

Ser Ser Arg Leu Ile His His Gln Arg Leu His His Gly Glu Lys

475

```
<210> 46
<211> 534
<212> PRT
<213> Homo sapiens
```

Lvs His

125

140

155

170

46/91

145

160

175

<220>

<221> misc-feature <223> Incyte ID No.: 3128913CD1

47/91

<400> 46 Met Ala Val Glu Ser Gly Val Ile Ser Thr Leu Ile Pro Gln Asp 15 10 Val Glu Asp Asn Phe Pro Pro Glu Gln Glu Leu Ile Leu Val Lys 25 ser Thr Gln Ser Cvs Ser Trp Asp Glu Lys Phe Lys Gln Asn Gly 40 35 Gln Glu Leu Phe Arg Gln Gln Phe Arg Lvs Tyr Gln Glu 60 50 Thr Pro Gly Pro Arg Glu Ala Leu Ser Arg Leu Gln Glu Leu 65 70 Pro Glu Leu His Thr Lys Glu Gln Ile Leu Tyr Gln Trp Leu Met 85 90 a n Glu Gln Phe Leu Ser Leu Pro Glu Glu Glu Leu Leu Val Leu Ile 95 100 Gln Gln His Asn Pro Glu Ser Gly Glu Glu Leu Gln Ile Trp Val 110 115 Glu Asp Leu Glu Arg Glu Phe Asp Asp Thr Leu Leu 125 130 135 Gln Gln Val Pro Ala Ser Pro Gln Gly Pro Ala Val Pro Trp 145 140 Lys Asp Leu Thr Cys Leu Arg Ala Ser Gln Glu Ser Tle 160 Lys Thr Gln Leu Lys CVS Leu Gln Pro Leu 180 170 175 Glu Thr Ala Thr Lvs Pro Lys Ser Asp Cys Glu Asn Ser 185 190 195 Pro Gln Glu Pro Glu Gly Ile Ser Glu Glu Lys Ser Gln Gly Leu 205 210 200 Val Trp Lys Phe Arg Gly Ile Ser Glu His Glu Ser 225 220 215 Gln Gly Ser Ala Thr Gly Glu Lys Leu Arg Ser Gln Gly 240 230 235 Phe Ser Gln Val Ile Phe Thr Asn 255 245 250 Thr Leu Ile Leu Thr Asp Leu Tyr Asp Glu Ala Glu Arg Cys 265 270 260 Gln Lys Val Pro Pro Glu Glu Arg Pro Tyr Ser Ile Met Cvs 275 280 285 Gly His Ser Phe Lys Leu Arg Cys Asp Val Cys 300 290 295 Cys Ile His Thr Gly Glu Lys Pro Tyr Lys Thr Gln His Gln Arg 315 305 310 Ala Phe Ser Leu Arg Ser Tyr Ile Asn Gln Cys Gly Lys 330 325 320 His Gln Arg Ile His Ser Gly Glu Lys Ala Tyr Glu Cys Ser Glu 345 335 340 Asn Gln Ser Ser Ala Leu Ile Arg His Cys Gly Lys Ala Phe 355 360 350 Glu Lys Ala Cys Lys Cys Asn Glu Cys Gly Ile His Thr Gly 365 370 375 Ile Ile Ile His Gln Arg Ala Phe Ser Gln 390 380 385 Pro Tyr Glu Cys Asn Glu Cys Gly Lys Thr His Thr Gly Glu Lys 395 405 400 Gln Arg Ile His Thr Ser Gln Ser Ser Lys Leu Ile Arg His 420 410 415 Gly Glu Arg Pro Tyr Glu Cys Asn Glu Cys Ala Phe Arg 435 425 430 Ile Thr His Gln Arg Ile His Ser Gly Glu Gln Ser Ser Glu Leu 450 440 445 Lys Pro Tyr Glu Cys Ser Glu Cys Gly Lys Ala Phe Ser Leu Ser

PCT/US00/02237

PCT/US00/02237

```
465
                455
                                      460
Ser Asn Leu Ile Arg His Gln Arg Ile His Ser Gly Glu Glu Pro
                                      475
                470
Tyr Gln Cys Asn Glu Cys Gly Lys Thr Phe Lys Arg Ser Ser Ala
                485
                                      400
Leu Val Gln His Gln Arg Ile His Ser Gly Asp Glu Ala
                                                          510
                500
                                     505
Cys Asn Glu Cys Gly Lys Ala Phe Arg His Arg Ser Val Leu Met
                                                          525
                515
                                     520
```

Arg His Gln Arg Val His Thr Ile Lys 530

```
<210> 47
<211> 206
<212> PRT
```

<213> Homo sapiens

<220> <221> misc-feature <223> Incyte ID No.: 3282941CD1

<400> 47 Met Ser Thr Gly Ser Val Ser Asp Pro Glu Glu Met Glu Leu Arg 15 5 10 1 Ser Lys Arg Pro Pro Gly Leu Gln Arg Glu Tyr Pro Val Pro Ala 25 20 Leu Arg Gly Val Glu Arg Ser Tyr Ala Ser Pro Ser Asp Asn Ser 45 35 40 Ser Ala Glu Glu Glu Asp Pro Asp Gly Glu Glu Glu Arg Cys Ala 50 55 60 Leu Gly Thr Ala Gly Ser Ala Glu Gly Cys Lys Arg Lys Arg Pro Arg Val Ala Gly Gly Gly Gly Ala Gly Gly Ser Ala Gly Gly Glv 90 85 20 Gly Lys Lys Pro Leu Pro Ala Lys Gly Ser Ala Ala Glu Cys 105 95 100 Gln Ser Gln Arg Asn Ala Ala Asn Ala Arg Glu Arg Ala Arg Met 110 115 120 Arg Val Leu Ser Lys Ala Phe Ser Arg Leu Lys Thr Ser Leu Pro 130 135 125 Thr Lys Leu Ser Lys Leu Asp Thr Leu Arg Trp Val Pro Pro Asp 145 150 140 Leu Ala Ser Ser Tyr Ile Ala His Leu Arg Gln Leu Leu Gln Glu 165 155 160 Asp Arg Tyr Glu Asn Gly Tyr Val His Pro Val Asn Leu Thr Trp 175 180 170 Pro Phe Val Val Ser Gly Arg Pro Asp Ser Asp Thr Lys Glu Val 185 190 195 Ser Ala Ala Asn Arg Leu Cys Gly Thr Thr Ala 200 205

<210> 48 <211> 172

<212> PRT <213> Homo sapiens

<220>

PCT/US00/02237

```
<221> misc-feature
<223> Incyte ID No.: 3286656CD1
Met Glu Ser Val Thr Phe Glu Asp Val Ala Val Glu Phe Ile Gln
                                      1 0
                                                           15
Glu Trp Ala Leu Leu Asp Ser Ala Arg Arg
                                         Ser Leu Cys Lys Tyr
                 20
                                      25
                                                           3 0
Arg Met Leu Asp Gln Cys Arg Thr Leu Ala
                                         Ser Arg Gly Thr Pro
                 35
                                      40
Pro Cys Lys Pro Ser Cys Val Ser Gln Leu Gly Gln Arg Ala Glu
                 50
                                      55
                                                           60
Pro Lys Ala Thr Glu Arg Gly Ile Leu Arg Ala Thr Gly Val Ala
                 65
                                      70
                                                           75
Trp Glu Ser Gln Leu Lys Pro Glu Glu Leu Pro Ser Met Gln Asp
                 80
                                      85
                                                           90
Leu Leu Glu Glu Ala Ser Ser Arg Asp Met Gln Met Gly Pro Gly
                 95
                                     100
                                                          105
Leu Phe Leu Ard Met Gln Leu Val Pro Ser
                                         Ile Glu Glu Arg
                                                          Glu
                                                          120
                110
                                     115
Thr Pro Leu Thr Arg Glu Asp Arg Pro Ala Leu Gln Glu Pro
                                                         Pro
                125
                                     130
                                                          135
Trp Ser Leu Gly Cys Thr Gly Leu Lys Ala Ala Met Gln Ile Gln
                140
                                     145
Arg Val Val Ile Pro Val Pro Thr Leu Gly His Arg Asn Pro Trp
                155
                                     160
                                                          165
Val Ala Arg Asp Ser Ala Met
                170
```

```
<210> 49
<211> 275
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 3490802CD1
```

<400> 49 Met Gly Pro Leu Gln Phe Arg Asp Val Ala Ile Glu Phe Ser Leu 10 15 Leu Asp Thr Ala Gln Arg Asn Leu Tyr Arg Glu Glu Trp His Cvs 20 25 3.0 Asp Val Met Leu Glu Asn Tyr Arg Asn Leu Val Phe Leu Gly 35 40 Val Val Ser Lys Pro Asp Leu Val Thr Cys Leu Glu Gln Gly Lys 55 60 50 Lys Pro Leu Thr Met Glu Arg His Glu Met Ile Ala Lys Pro Pro 65 Val Met Ser Ser His Phe Ala Gln Asp Leu Trp Pro Glu Asn Ile 80 Gln Asn Ser Phe Gln Ile Gly Met Leu Arg Arg Tyr Glu Glu Cys 100 105 95 Ard His Asp Asn Leu Gln Leu Lys Lys Gly Cys Lys Ser Val Gly 115 120 Glu His Lys Val His Lys Gly Gly Tyr Asn Gly Leu Asn Gln Cys 125 130 Leu Thr Thr Gln Lys Glu Ile Phe Gln Cys Asp Lys Tyr Gly

PCT/US00/02237

WO 00/44900

140 145 Phe Ser Asn Ser Asn Thr Tyr Lys Thr Arg Lvs Val Phe His Lvs 160 155 His Thr Gly Ile Asn Leu Phe Lys Cys Ile Ile Cys Gly Lys Ala 175 170 Thr Phe Lys Arg Ser Ser Thr Leu Thr Thr His Lys Lys 190 185 Gly Lys Ala Phe Asn Gly Glu Lys Pro Tyr Lys Cys Glu Glu Cys 205 210 200 Ile His Thr Gly Glu Gin Ser Ser Asn Leu Thr Thr His Lys Arg 220 225 215 Cys Glu Glu Cys Gly Lys Ala Phe Asn Trp Ser Lvs Pro Tvr Lvs 240 230 235 Ile Glu Arg Lys Ser Asp Leu Asn Lys His Lys Lys Ile His 255 245 250 Tyr Ile Val Lys Asn Val Thr Asp Leu Leu Asn Val Pro Pro Leu 265 260 Leu Ile Ser Ile Arg 275

<210> 50 <211> 236

<212> PRT <213> Homo sapiens

<220>

<221> misc-feature <223> Incyte ID No.: 3507366CD1

<400> 50 Met Asp Lys Arg Tyr Leu Gln Phe Asp Ile Lys Ala Phe Val Glu 10 15 5 Ile Lys Trp Cys Pro Thr Pro Gly Cys Asp Arg 30 20 25 Asp Thr Ser Gly Ser Ala Val Arg Leu Thr Lys Gln Gly Ser Asn 45 40 35 Thr Leu Ser Phe Pro Leu Leu Arg Ala Pro Ala Val Asp Cys Gly 55 60 50 Lys Gly His Leu Phe Cys Trp Glu Cys Leu Gly Glu Ala His Glu 70 75 65 Leu Gln Lys Ile Thr Pro Cys Asp Cys Gln Thr Trp Lys Asn Trp 90 85 80 Glu Glu Met Lys Pro Glu Glu Leu Val Gly Val Ser Glu Ala Tyr 100 105 95 Cys Asp Ala Ala Asn Cys Leu Trp Leu Leu Thr Asn Ser Lys Pro 115 110 Ala Asn Cys Lys Ser Pro Ile Gln Lys Asn Glu Gly Cys Asn His 135 125 130 Cys Lys Tyr Asp Phe Cys Trp Ile Cys Leu Met Gln Cys Ala Lys 150 145 140 Glu Glu Trp Lys Lys His Ser Ser Ser Thr Gly Gly Tyr Tyr Arg 165 160 155 Cys Thr Arg Tyr Glu Val Ile Gln His Val Glu Glu Gln Ser Lys 180 175 170 Glu Met Thr Val Glu Ala Glu Lys Lys His Lys Arg Phe Gln Glu 195 190 185 Leu Asp Arg Phe Met His Tyr Tyr Thr Arg Phe Lys Asn His Glu 205 210 200 His Ser Tyr Gln Leu Glu Gln Arg Leu Leu Lys Thr Ala Lys Glu

<400> 52

PCT/US00/02237

```
225
                215
                                     220
Lys Met Glu Cln Met Ser Arg Val Ser Lys Asn
                230
                                     235
<210> 51
<211> 214
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 3573060CD1
-400× 51
Met Asn Leu Ser Ser Ala Ser Ser Thr Glu Glu Lys Ala Val Thr
                                                           15
 1
                  5
                                      10
Thr Val Leu Trp Gly Cys Glu Leu Ser Gln Glu Arg Arg Thr Trp
                                                           30
                 20
Thr Phe Arg Pro Gln Leu Glu Gly Lys Gln Ser Cys Arg Leu Leu
                 35
                                      40
Leu His Thr Ile Cys Leu Gly Glu Lys Ala Lys Glu Glu Met His
                 50
                                      55
                                                           60
Arg Val Glu Ile Leu Pro Pro Ala Asn Gln Glu Asp Lvs Lvs Met
                 65
                                      70
Gln Pro Val Thr Ile Ala Ser Leu Gln Ala Ser Val Leu Pro Met
                                      25
                 90
Val Ser Met Val Gly Val Gln Leu Ser Pro Pro Val Thr Phe Gln
                 95
                                     100
                                                          105
Leu Arg Ala Gly Ser Gly Pro Val Phe Leu Ser Gly Gln Glu Arg
                                                          120
                110
                                     115
Tyr Glu Ala Ser Asp Leu Thr Trp Glu Glu Glu Glu Glu Glu Glu
                125
                                     130
Gly Glu Glu Glu Glu Glu Glu Glu Asp Asp Glu Asp Glu Asp
                                                          150
                140
                                    145
Ala Asp Ile Ser Leu Glu Glu Gln Ser Pro Val Lys Gln Val Lys
                155
                                    160
                                                          165
Arg Leu Val Pro Gln Lys Gln Ala Ser Val Ala Lys Lys Lys Lys
                170
                                    175
                                                          180
Leu Glu Lys Glu Glu Glu Glu Ile Arg Ala Ser Val Arg Asp
                                                         Lvs
                185
                                    190
                                                          195
Ser Pro Val Lys Lys Ala Lys Ala Thr Ala Arg Ala Lys Lys Pro
                200
                                    205
                                                         210
Glv Phe Lys Lys
<210> 52
<211> 396
<212> PRT
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 3573661CD1
```

51/91

Met Asn Phe Thr Val Gly Phe Lys Pro Leu Leu Gly Asp Ala His

and the state of t

PCT/US00/02237

1				5					10					15
Ser	Met	Asp	Asn	Leu 20	Glu	Lys	Gln	Leu	11e 25		Pro	Ile	Cys	: Lei
Glu	Met	Phe	Ser	Lys 35	Pro	Va1	Va1	Ile	Leu 40		Cys	Gln	His	Asr 49
Leu	Cys	Arg	Lys	Cys 50	Ala	Asn	Asp	Val	Phe 55		Ala	Ser	Asn	Pro 60
Leu	Trp	Gln	Ser	Arg 65	G1y	Ser	Thr	Thr	Val		Ser	Gly	Gly	Arg
Phe	Arg	Cys	Pro		Cys	Arg	His	Glu		Val	Leu	Asp	Arg	
Gly	Val	Tyr	Gly		Gln	Arg	Asn	Val		Val	Glu	Asn	Ile	
Asp	Ile	Tyr	Lys		Glu	Ser	Ser	Lys			His	Ser	Lys	
Glu	Gln	His	Leu		Cys	Glu	Glu	His		Glu	Glu	Lys	Ile	
Ile	Tyr	Суѕ	Leu		Cys	Glu	Val	Pro		Cys	ser	Leu	Суѕ	
Val	Phe	Gly	Ala		Lys	Asp	Суз	Glu			Pro	Leu	Pro	
Ile	Tyr	Lys	Arg		Lys	Ser	Glu	Leu			Gly	Ile	Ala	
Leu	Val	Ala	Gly		Asp	Arg	Val	Gln		Val	Ile	Thr	Gln	
Glu	Glu	Val	Cys		Thr	Ile	Glu	Asp		Ser	Arg	Arg	Gln	
Gln	Leu	Leu	Asn		Arg	Phe	Glu	Ser		Cys	Ala	Val	Leu	
Glu	Arg	Lys	Gly		Leu	Leu	Gln	Ala		Ala	Arg	Glu	Gln	
Glu	Lys	Leu	Gln		Val	Arg	Gly	Leu		Arg	Gln	Tyr	Gly	
His	Leu	Glu	Ala	Ser 260	Ser	Lys	Leu	Val	Glu 265	Ser	Ala	Ile	Gln	Ser 270
Met	Glu	Glu	Pro		Met	Ala	Leu	Tyr		Gln	Gln	Ala	Lys	Glu 285
Leu	Ile	Asn	Lys	Val 290	Gly	Ala	Met	Ser	Lys 295	Val	Glu	Leu	Ala	Gly 300
Arg	Pro	Glu	Pro		Tyr	Glu	Ser	Met		Gln	Phe	Thr	Val	Arg 315
Va1	Glu	His	Val	Ala 320	Glu	Met	Leu	Arg	Thr 325	Ile	Asp	Phe	Gln	Pro 330
Gly	Ala	Ser	Gly		Gly	Arg	Gly	Gly	Gly 340	Pro	Arg	Arg	Lys	Lys 345
Arg	Ala	Thr	Arg		Pro	Glu	Glu	Lys		Ala	Arg	Met	Gly	
Tyr	Arg	Pro	Leu	Arg 365	Pro	Asn	Pro	Asp	Pro 370	Leu	Leu	Arg	Lys	Ser 375
Pro	Arg	Arg	Leu		Ile	Ser	Gly	Gly	Arg 385	Asn	Ser	Cys		Lys 390
Lys	Thr	Pro	Ala		Phe									

<210> 53 <211> 486 <212> PRT <213> Homo sapiens

<221> misc-feature

<220>

PCT/US00/02237

<223> Incvte ID No.: 3633422CD1 Met Arg Arg Leu Val His Asp Leu Leu Pro Pro Glu Val Cys Ser 10 15 7 5 Asn Glu Ile Ser Leu Leu Asn Pro Ala Ala Ile Tyr Ala Asn Leu 20 25 3 0 Val Glu Val Tyr Asp Tyr Thr Leu Tyr Gly Phe Asp 35 40 15 His Pro Glu Ile Phe Ser Thr Ala Arg Ala Asp Ala 50 55 60 Tyr Pro Leu Ile Glu His Tvr Lvs Glu Gly Ile Arg Lys 75 70 65 Ser Phe Ala Ile Arg Gly Leu His Tyr Asp Tyr Asn Pro 90 80 85 Leu Met Lys Ile Asp Val Lvs Ser Leu Ala Phe His Tvr 95 100 105 Glu Tyr Arg Gly Leu Gln Pro Val Pro Asp Gln Leu Glv Thr Ala 115 120 110 Tyr Gly Gly Thr Gln Tyr Glu Val Ile Glu Leu 125 135 Tyr Gly Lys Gly Pro Phe Gln Met Ser Gly Phe 150 140 145 Leu Pro Glu Met Ala Asp Ile Phe Ser 160 165 155 Tyr Phe Leu Gly His Ser Leu Glu Phe Asp Gln Ala His 175 180 170 Lvs Leu Tvr Lvs Asp Val Thr Asp Ala Ile Arg 190 195 185 Trp Ile Glu Gln Asp Ile Gly Leu Met Tyr Gln 200 205 210 Thr Phe Ala Val Leu Ser Arg Leu Val Arg Gly Asp Glu 225 215 220 Phe Gly Lys Gln Leu Phe Leu Ile Thr Asn 235 240 230 Val Gly Gln Asp Lys Gly Met 250 255 245 Val Ile Val Gln Ala Asp Phe Leu Phe Asp Val 265 270 260 Asp Glu Lys Gly Ser Thr Asp Arg Arg Lys Pro Phe Arg Lys Leu 285 275 280 Ile Thr Arg Leu Glu Lvs Glv Lvs Leu Gln Trp Asp Arg 290 295 300 Arg Gln Gly Asn Leu Phe Asp Phe Arg Leu Arg 315 305 310 Tyr Phe Gly Asp His Gly Pro Arg Val Leu 330 320 325 Thr Gly Ala Ile Ile Ala Asp Leu Met Leu His Gly Trp Arg 345 335 340 Pro Glu Leu Glu Arg Glu Ile Arg Ile Ile Tvr 355 360 350 His Ser Leu Thr Trp Gln Gln Ala Leu Glu 375 365 370 Arg Gln Val Leu Ala Arg Met Gln Thr Gln Asp Ala Glu Ser 390 385 380 Ala Trp Met Lys Glu Arg Gln Glu Leu Arg Ala 400 395 Leu Phe Asn Ala Gln Phe Gly Ser Ile Phe 410 415 420 Pro Thr Tyr Phe Ser Arg Arg Leu Val Arg Phe Ser Asp Leu Tyr

. Jan 10 60

Laborate Colors, Line of

PCT/US00/02237

```
425
                                                           435
                                      430
Met Ala Ser Leu Ser Cys Leu Leu Asn Tyr Arg Val Asp Phe Thr
                                                           450
                 440
Phe Tyr Pro Arg Arg Thr Pro Leu Gln His Glu Ala Pro Leu
                                                           Trp
                 155
                                      460
                                                           465
Met Asp Gln Leu Cys Thr Gly Cys Met Lys Thr Pro Phe Leu Gly
                                                           480
                 470
                                      475
Asp Met Ala His
                Ile Arg
                 485
```

<210> 54 <211> 555 <212> PRT

<213> Homo sapiens

<220>

<221> misc-feature

<223> Incyte ID No.: 3993377CD1

Met Gly Ala Glu Asp Lys Leu Pro Leu Glu Asp Ser Pro Val Ile 5 10 Ala Ala Leu Asp Cys Pro Ser Leu Asn Asn Ala Thr Ala Phe Ser 20 25 30 ser Leu Leu Ala Asp Asp Ser Gln Thr Ser Thr Ser Ile Phe Ala 35 40 45 Pro Thr Ser Pro Pro Val Leu Gly Glu Ser Val Leu Gln Asp Asn 50 55 60 Glu Gln Glu Glu Met Ser Phe Asp Leu Asn Asn Gly Ser Asp Ala 70 75 65 Glu Thr Gln Ser Ser Asp Phe Pro Pro Ser Leu Thr Gln Pro Ala 80 85 90 Pro Asp Gln Ser Ser Thr Ile Gln Leu His Pro Ala Thr Ser Pro 95 100 105 Thr Ser Pro Ala Val Ser Leu Val Val Ser Ala Val Ser Pro Thr 115 120 110 Pro Ala Ala Ser Pro Glu Ile Ser Pro Glu Val Cys Pro Ala Ala 125 130 Ser Thr Val Val Ser Pro Ala Val Phe Ser Val Val Ser Pro Ala 150 145 140 Ser Ser Ala Val Leu Pro Ala Val Ser Leu Glu Val Pro Leu 155 160 165 Ala Ser Val Thr Ser Pro Lys Ala Ser Pro Val Thr Ser Pro Ala 170 175 Ala Ala Phe Pro Thr Ala Ser Pro Ala Asn Lys Asp Val Ser Ser 185 190 Gly Phe Leu Glu Thr Thr Ala Asp Val Glu Glu Ile Thr Gly Glu 200 205 210 Leu Thr Ala Ser Gly Ser Gly Asp Val Met Arg Arg Arg Ile Ala 215 220 225 Arg Leu Pro Leu Gln His Gly Trp Arg Thr Pro Glu Glu Val Ara 230 235 240 Glu Val Arg Ile Lvs Asn Ser Ser His Arg Trp Gln Gly Glu Thr 245 250 255 Trp Tyr Tyr Gly Pro Cys Gly Lys Arg Met Lys Gln Phe Pro Glu 260 265 270 Val Ile Lys Tyr Leu Ser Arg Asn Val Val His Ser Val Arg Ara 285 275 280 Glu His Phe Ser Phe Ser Pro Arg Met Pro Val Gly Asp Phe Phe

2 1 to 2 2 10

PCT/US00/02237

```
200
                290
                                     295
Glu Glu Arg Asp Thr Pro Glu Gly Leu Gln Trp Val Gln Leu Ser
                                                           315
                305
                                     310
Ala Glu Glu Ile Pro
                    Ser Arg Ile Gln Ala Ile Thr Gly Lys Arg
                                     325
                                                           330
                320
Gly Arg Pro Arg Asn Thr Glu Lys Ala Lys Thr Lys Glu Val
                                                          Pro
                                                           345
                335
                                     340
   Val Lys Arg Gly Arg Gly Arg Pro Pro Lys Val Lys Ile
                                                          Thr
                350
                                     355
                                                          360
Glu Leu Leu Asn Lys Thr Asp Asn Arg Pro Leu Lys Lys Leu Glu
                365
                                     370
                                                          375
Ala Gln Glu Thr Leu Asn Glu Glu Asp Lys Ala Lys Ile Ala Lys
                                     385
                                                          390
                380
Ser Lvs Lvs Lvs Met Arg Gln Lvs Val Gln Arg Glv Glu Cvs Gln
                395
                                     400
                                                          405
Thr Thr Ile Gln Gly Gln Ala Arg Asn Lys Arg Lys Gln Glu Thr
                410
                                     415
                                                          420
                                        Lys Ser Lys Ala Glu
Lys Ser Leu Lys Gln Lys Glu Ala Lys Lys
                425
                                     430
                                                          435
Lys Glu Lys Gly Lys Thr Lys Gln Glu Lys
                                         Leu Lys Glu Lys
                                                          Va1
                440
                                     445
                                                          450
Lys Arg Glu Lys Lys Glu Lys Val Lys Met Lys Glu Lys Glu Glu
                155
                                     460
                                                          465
   Thr Lys Ala Lys Pro Ala Cys Lys Ala Asp Lys Thr Leu Ala
                                     475
                                                          480
                470
Thr Gln Arg Arg Leu Glu Glu Arg Gln Arg Gln Gln Met Ile
                                                          Leu
                                     490
                                                          495
                485
                                                          His
Glu Asp Met Lys Lys Pro Thr Glu Asp Met Cys Leu Thr Asp
                500
                                     505
                                                          510
Gln Pro Leu Pro Asp Phe Ser Arg Val Pro Gly Leu Thr Leu
                                                          Pro
                515
                                     520
                                                          525
Ser Gly Ala Phe Ser Asp Cys Leu Thr Ile Val Glu Phe Leu
                                                          His
                                                          540
                530
                                     535
Ser Phe Gly Lys Val Leu Gly Leu Asp Pro Ala Gln Gly Cys
                                                          Ala
                                                          555
                545
                                    550
```

Lys Met Gly Thr Gly Trp Glu Gly Phe Gln Arg Thr Leu Lys Glu 35 40 40 Val Ala Tyr Ile Leu Leu Cys Cys Trp Cys Ile Lys Glu Leu Leu 50 55 Asp

<210> 56

PCT/US00/02237

```
<211> 2781
<212> DNA
<213> Homo sapiens
<220×
<221> misc-feature
<223> Incyte ID No.: 025733CB1
gaagtggggt gcacgcttcg ggttggtgtc atggcagctg cggggagccg caagaggcgc 60
ctggeggage tgaeggtgga cgagtteeta getteggget ttgaeteega gteegaatee 120
gagteegaaa atteteeaca ageggagaca egggaageac gegaggetge eeggagteeg 180
gataagccgg gegggagccc ctcggccagc cggcgtaaag gccgtgcctc tgagcacaaa 240
gaccagetet eteggetgaa ggacagagac eeegagttet acaagtteet geaggagaat 300
gaccagagec tgetaaactt cagegacteg gacagetetg aggaggaaga ggggeegtte 360
cactecetge cagatgtget ggaggaagee agtgaggagg aggatggage ggaggaagga 420
gaagatgggg acagagteee cagagggetg aaggggaaga agaattetgt teetgtgaee 480
gtcgccatgg ttgagagatg gaagcaggca gcaaagcaac gcctcactcc aaagctgttc 540
catgaagtgg tacaggcgtt ccgagcagct gtggccacca cccgagggga ccaggaaagt 600
gctgaggcca acaaattcca ggtcacggac agtgctgcat tcaatgctct ggttaccttc 660
tgcatcagag acctcattgg ctgtctccag aagctgctgt ttggaaaggt ggcaaaggat 720
agcagcagga tgctgcagcc gtccagcagc ccgctctggg ggaagetteg tgtggacatc 780
aaggettace tgggetegge catacagetg gtgteetgte tgteggagae gaeggtgttg 840
geggeegtge tgeggeacat cagegtgetg gtgeeetget teetgaeett ceecaagcag 900
tgccgcatgc tgctcaagag aatggtggtc gtatggagca ctggggagga gtctctgcgg 960
gtgetggett teetggteet eageagagte tgeeggeaca agaaggacae ttteettgge 1020
cccgtcctca agcaaatgta catcacgtat gtgaggaact gcaagttcac ctcgcctggt 1080
geoeteecet teateagett catgeagtgg acettgaegg agetgetgge cetggageeg 1140
ggtgtggcct accagcacgc cttcctctac atccgccagc tcgccataca cctgcgcaac 1200
gccatgacca cccgcaagaa ggaaacatac cagtctgtgt acaactggca gtatgtgcac 1260
tgeetettee tgtggtgeeg ggteetgage aetgegggee ceagegaage cetecageee 1320
ttggtctacc cccttgccca agtcatcatt ggctgtatca agctcatccc cactgcccgc 1380
ttetaccege tgegaatgea etgeateegt geeetgacge tgeteteggg gagetegggg 1440
geetteatee eggtgetgee ttteateetg gagatgttee ageaggtega etteaacagg 1500
aagccagggc gcatgagctc caagcccatc aacttctccg tgatcctgaa gctgtccaat 1560
gtcaacctgc aggagaaggc gtaccgggac ggcctggtgg agcagctgta cgacctcacc 1620
ctggagtace tgcacageca ggcacaetge ateggettee eggagetggt getgeetgtg 1680
gteetgeage tgaagtegtt cetcegggag tgeaaggtgg ceaactactg ceggeaggtg 1740
cagcagetge tigggaaggt teaggagaac teggeataca tetgeageeg eegecagagg 1800
gttteetteg gegtetetga geageaggea gtggaageet gggagaaget gaecegggaa 1860
gaggggacac cettgacett gtactacage cactggegea agetgegtga eegggagate 1920
cagetggaga teagtggeaa agageggetg gaagaeetga aetteeetga gateaaacga 1980
aggaagatgg ctgacaggaa ggatgaggac aggaagcaat ttaaagacct ctttgacctg 2040
aacagetetg aagaggacga cacegaggga tteteggaga gagggatact gaggeceetg 2100
agcactegge atggggtgga agacgatgaa gaggacgagg aggagggega ggaggacage 2160
agcaactcgg agggtgaatg gtcttgggat ggagacccag acgcagaggc ggggctggcc 2220
cetggggage tgcagcaget ggcccagggg ceggaggacg agetggagga tetgcagete 2280
tcagaggacg actgaggcag cccatctggg gggcctgtag gggctgccgg gctggtggcc 2340
agtgtttcca cctccctggc agtcaggcct agaggctggc gtctgtgcag ttgggggagg 2400
cagtagacac gggacagget ttattattta tttttcagca tgaaagacca aacgtatcga 2460
gagetggget gggetggget ggtgtggetg etgaageeee acagetgtgg getgetgaag 2520
teageteege gggggagetg accetgaegt cagcagaeeg agaccagtee cagttecagg 2580
gggaggcctg caggcccctg gccccttcca ccacctctgc cctccgtctg cagacctcgt 2640
ccatctgcac caggetetge etteactece ccaagtettt gaaaatttgt teettteett 2700
tgaagteaca ttttctttta aaattttttg ttttgcatcc gaaaccgaaa gaaataaagc 2760
                                                                   2781
ggtgggaggc aaaaaaaaa a
```

<210> 57 <211> 2544

<212> DNA

CLEANED AND SHEET AND A PROPERTY OF THE PROPERTY OF THE PARTY OF THE P

WO 00/44900 <213> Homo sapiens

PCT/US00/02237

```
<220>
<221> misc-feature
<223> Incyte ID No.: 079702CB1
<400> 57
cgggaaacca aaatggcgag qqqctgtatt gaagtgggct gtgtttgagg ccggtgtaag 60
aacgeteatt etacceccaa ceettgtete caaggacete ggtttgtgeg tgcatatgtg 120
cogggtacce ggtggggcgg gtgcccagta agtgctcgga ctcgcagggg aagcgccac 180
ggggacggat tggttgtttt ttcctgtatg aagcggttgg caccactgaa gtgaccgaat 240
gagagactet acaggggcag gtaatteact ggtecacaag eggteteett taegtegaaa 300
ccaaaagacc ccaacatcct tgaccaagct gtctttacag qatggacata aagccaaaaa 360
gccagcatgt aaatttgaag agggtcagga tgtcctagct agatggtcag atggcttgtt 420
ttatottgge actatoaaaa agataaacat attgaaacag agotgettea teatattga 480
agacagttet aaateetggg ttetetggaa ggacatteaa acaggageea etggaagtgg 540
ggaaatggtc tgtacaatat gtcaagaaga gtattcagaa gctcccaatg aaatggttat 600
atgtgacaag tgtggccaag gatatcatca gttgtgtcac acacctcata ttgattccag 660
tgtgattgat tcagatgaaa aatggctctg tcggcagtgt gtttttgcaa caacaacaaa 720
gaggggtggt gcacttaaga aaggaccaaa tgccaaagca ttgcaagtca tgaagcagac 780
attaccetat agtgtggcag acettgaatg ggatgcaggt cataaaacca atgtccagca 840
gtgttactgc tattgtggag gccctggaga ctggtatttg aagatgctac agtgctgcaa 900
atgtaagcag tggtttcatg aggcttgtgt gcaatgcctt caaaagccaa tgctatttgg 960
agacagattt tatacgttta tatgctctgt ctgcagttct ggaccagaat acctcaaacg 1020
totaccatta cagtgggtag atatagcaca cotatgcott tacaacctaa gtgttattca 1080
taaqaaqaaa tactttgatt ctgaacttga gcttatgaca tacattaatg aaaactggga 1140
tagattgcac cetggagage tggcagacac accaaaatct gaaagatatg agcatgttct 1200
ggaggcatta aatgattaca agaccatgtt tatgtctqqq aaaqaaataa aqaaqaagaa 1260
gcatttgttt gggttgcgaa ttcgtgttcc tcctgtgcca ccaaatgtgg ctttcaaagc 1320
agagaaaqaa cctgaaggaa catctcatga atttaaaatt aaaggcagaa aggcatccaa 1380
acctatatet gatteaaggg aagtaageaa tggcatagaa aaaaaaaaaa aaaaaaaate 1440
tgtaggtcgt ccacctggcc catatacaag aaaaatgatt caaaaaactg ctgagccact 1500
tttggataag gaatcaattt cagagaatcc tactttggat ttaccttgtt ctatagggag 1560
aactgaggga actgcacatt catccaatac ctcagatgtg gatttcacgg gtgcttccag 1620
tgcaaaagaa actactcgt ctagcattte caggcattat ggattatetg actccagaaa 1680
aagaacgcgt acaggaagat cttggcctgc tgcaatacca catttgcgga gaagaagagg 1740
tegtetteca agaagageae tecagactea gaacteagaa attgtaaaaq atgatgaaqq 1800
caaagaagat tatcagtttg atgaactcaa cacagagatt ctgaataact tagcagatca 1860
ggagttacaa ctcaatcatc taaagaactc cattaccagt tattttggtg ctgcaggtag 1920
aatagcatgt ggcgaaaaat accgagtttt ggcacgtcgg gtgacacttg atggaaaggt 1980
gcagtatett gtggaatggg aaggageaac tgcateetga etgtaggaet gaacattatg 2040
ttcactgcac totgattttc tqtaggtaca gttcaaagcc ctaaaggagt ctggctttta 2100
ctatetttet taaaaaaaaa aaaaagteaa aaaaatteaa aaaaggggat gataetagee 2160
ttaacatgta cctgtcaatg ttatggatat tgtcataaaa aggtatcttt taaaaatcag 2220
aacagagact taattittta aatcttaaga titgtagaat gittctagga taggatatta 2280 aaaatgattg aaacccatgc atggtgttag acaattittc taattattcc attgagtcag 2340
ttttttgtga ttagtgatta tcagagcaaa catcatgtag atagcacaag tatttggaga 2400
aacgttgttt gttttgttac caaaatgttg gaaaaattta tttcaatacc ttttagattt 2460
cataaagtgc agtgtatata atgcctactg aaagactgta aaatattgaa attttctttc 2520
aagcaaagtg taaataaata tato
<210> 58
<211> 1627
<212> DNA
```

<213> Homo sapiens

<220> <221> misc-feature

<223> Incyte ID No.: 116208CB1

The state of the s

<400> 58

PCT/US00/02237

```
ctgatgaggg cgctgcattt attgaagagc ggctgcagcc ctgcggttca gattaaaatc 60
cgagaattgt atagacgccg atatccacga actottgaag gactttctga tttatccaca 120
atcaaatcat cggttttcag tttggatggt ggctcatcac ctgtagaacc tgacttggcc 180
gtggetggaa tecactegtt geetteeact teagttacae etcacteace atcetetect 240
gttggttctg tgctgcttca agatactaag cccacatttg agatgcagca gccatctccc 300
ccaatteete etgtecatee tgatgtgeag ttaaaaaate tgecetttta tgatgteett 360
gatgttctca tcaagcccac gagtttagtt caaagcagta ttcagcgatt tcaagagaag 420
ttttttattt ttgctttgac acctcaacaa gttagagaga tatgcatatc cagggatttt 480
ttgccaggtg gtaggagaga ttatacagtc caagttcagt tgagactttg cctggcagag 540
acaagttgcc ctcaagaaga taactatcca aatagtctat gtataaaagt aaatgggaag 600
ctattteett tgeetggeta tgeaceaeeg cetaaaaatg ggattgaaca gaagegeeet 660
ggacgcccct tgaatattac atctttagtt aggttatctt cagctgtgcc aaaccaaatt 720
tocatttott gggcatcaga aattgggaag aattactota tgtotgtata tottgtacgg 780
cagettacat cagecatgtt attacagaga ttaaaaatga aaggtattag aaaccetgat 840
cattocagag cactaagtaa agaaaaactt actgcagatc ctgatagtga aattgctaca 900
actageette gggtateett gatgtgeest ttaggaaaaa tgaggetgac aateecatge 960
cottocagtoa ettotacaca tetocagtot tttgatocto coetetatet acaaatgaat 1020
gagaaaaagc ccacctggat ttgtcctgtg tgtgacaaaa aagctgccta tgaaagtcta 1080
atattagatg ggettttat ggaaattete aatgaetgtt etgatgtaga tgagatcaaa 1140
ttccaagaag atggttcttg gtgtccaatg agaccgaaga aagaagctat gaaagtatcc 1200
agccaaccqt qtacaaaaat agaaagttca agcgtcctca gtaagccttg ttcagtgact 1260
gtagccagtg aggcaagcaa gaagaaagta gatgttattg atcttacaat agaaagctct 1320
tetgacgaag aggaagaece teetgecaaa aggaaatgca tetttatgte agaaacacaa 1380
agcagcccaa ccaaaggggt totcatgtat cagccatott ctgtaagggt gcccagtgtg 1440
actteggttg atcetgetge tatteegeet teattaacag actacteagt accattecae 1500
catacgccaa tatcaagcat gtcatcagat ttgccaggag aacaaagaag aaatgatatt 1560
aataatgaac tgaagcttgg aacatcttot gatactgtgc aacagtgaat acaaaataaa 1620
                                                                  1627
accgata
<210> 59
<211> 1043
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incyte ID No.: 179261CB1
```

```
gtgggatatt tcagtcaaat gaaaatcatc tctgaaaatg tgcccagtta caaaactcat 60
gaatototta otttacotog gagaactoat gacagtgaga agcoctatga atacaaggaa 120
tatgagaagg tetteagttg tgaettagag tttgatgaat atcagaaaat acatactggt 180
ggaaaaaact atgaatgtaa tcaatgttgg aaaacctttg ggatagataa ctccagtatg 240
ttacaactga atattcatac tggtgtgaaa ccttgtaaat atatggaata tgggaataca 300
tgtagttttt ataaagactt taatgtatac cagaaaattc ataatgagaa qttctataaa 360
tgtaaggaat acagaaggac ctttgaaaga gttggaaaag ttactccact tcaaagagtt 420
catgatggtg agaaacactt tgaatgctca ttctgtggga aatcctttag agtgcatgca 480
caacttactc gacatcagaa aatccatact gatgagaaaa ettacaaatg tatggaatgt 540
ggcaaggact toagatttoa ttoacagott accgaacato agagaattoa tactggtgag 600
aaaccctaca aatgtatgca ctgtgagaag gtttttagaa ttagttcaca gctcattgaa 660
catcagagaa ttcacactgg tgagaaacct tatgcatgta aggaatgtgg gaaggctttt 720
ggagtatgta gagaacttgc tegtcatcag agaattcata ctggaaaata ctgtggatgg 780
atttaatagg taatcaaggc aattcagtat etecetetet taaagtetgt ttttagaett 840
catggtcatt ctgtatgtag acgtagaatt gcttagtcat agctgatata tacttatagc 900
tttgttagat gttgccaaat atttctcctt tttatgttaa aaagtttttt tcatgagttt 960
ctcatcctgg catgetttgt ttacaatagc ttttgatgtt tgtattattg ctcttttgac 1020
taatcagttt tttaattctg gat
```

in the surface of a secretary and was been properly by the property of the secretary of the

<210> 60 <211> 2448

<212> DNA <213> Homo sapiens

```
<220>
<221> misc-feature
<223> Incvte ID No.: 259161CB1
<400> 60
ctggcgggaa gattttactc ccgagtagcg gaaagatctg ctcgaggcct gggtgctttg 60
gtgtcggaga tccgagagtc ggagatcqqa qagtcggaca caqqacaqtc qqacaccqqa 120
cagtcaaaca coggagagtt agactgggct totoggtggg gagaggctot gggataacta 180
ctgttacage tttgaagggt caagggagga ttggccacca aagectgttt attagcaget 240
qccatttqtt qaaaqaaatt tqqattattt tagaaacaaa tttggaaaqa aaaagaatgg 300
cgtccgtttc agctctaact gaggaactgg attctataac cagtgagcta catgcagtag 360
aaattcaaat tcaagaactt acggaaaggc aacaagagct tattcagaaa aaaaaagtcc 420
tgacaaagaa aataaaggag tgtttagagg attctgatgc cggggcaagc aatgaatatg 480
attetteace tgeegettgg aataaagaag atttteeatg gtetggtaaa gttaaagata 540
ttctgcaaaa tgtctttaaa ctggaaaagt tcagaccact tcagcttgaa actattaacg 600
taacaatggc tggaaaggag gtatttettg ttatgcctac aggaggtgga aagagcttat 660
gttaccagtt accagcatta tgttcagatg gttttacact cgtcatttqc ccattgatct 720
ctcttatgga agaccaatta atggttttaa aacaattagg aatttcagca accatgttaa 780
atgettetag ttetaaggag catgttaaat gggtteatge tgaaatggta aataaaaact 840
ccgagttaaa gctgatttat gtgactccag agaaaattgc aaaaagcaaa atgtttatgt 900
caagactaga gaaagcctat gaagcaagga gatttactcg aattgctgtg gatgaagttc 960
actgctgtag tcagtgggga catgatttca gacctgatta taaggcactt ggtatcttaa 1020
agoggoagtt coctaacqca toactaattg ggotgactgc aactgcaaca aatcacgttt 1080
tgacggatgc tcagaaaatt ttgtgcattg aaaagtgttt tacttttaca gcttctttta 1140
ataagccaga tgtgaggttt gttatccatc attcaatgag taaatccatg gaaaattatt 1200
accaagagag tggacgtgca ggtcgagatg acatgaaagc agactgtatt ttgtactacg 1260
gctttggaga tatattcaga ataagttcaa tggtggtgat ggaaaatgtg ggacagcaga 1320
agetttatga gatggtatca tactgtcaaa acataagcaa atgtcgtcgt gtgttgatgg 1380
ctcaacattt tgatgaagta tggaactcag aagcatgtaa caaaatgtgc gataactgct 1440
gtaaagacag tgcatttgaa agaaagaaca taacagagta ctgcagagat ctaatcaaga 1500
teetgaagea ggeagaggaa etgaatgaaa aaeteaetee attgaaaetg attgattett 1560
ggatgggaaa gggtgcagca aaactgagag tagcaggtgt tgtggctccc acacttcctc 1620
gtgaagatet ggagaagatt attgcacact ttctaataca gcagtatett aaagaagact 1680
acagttttac agcttatget accatttegt atttgaaaat aggacetaaa getaatette 1740
tgaacaatga ggcacatgct attactatgc aagtgacaaa gtccacgcag aactctttca 1800
gggetgaate gteteaaaet tgteattetg aacaaggtga taaaaagatt ggaggaaaaa 1860
aattccaggc aacttccaga agaaggctgc aaacatgctt cagcaatctg gttctaagaa 1920
tacaggaget aagaaaagaa aaategatga tgeetgatat gaetgttact aaatttteta 1980
attaaagatg gtttatgcat gtatatgcca ttatttttgt agttagacaa tagtttttaa 2040
aagaatttoa tagatatttt atatgtatgg atotatattt toagagotta tototgaaga 2100
totaaacttt tgagaatgtt tgaaaattag agatcatgaa ttatataatt ttocagtata 2160
aaacaaggga aaaattttta tgtaaaaccc tttaaatgta aaatatttga gaataagttc 2220
atacaatcgt cttaagtttt ttatgccttt atatacttag ctatattttt tcttttgaca 2280
taactatett tttgaaagea atattataet gacagagget caetgagtga taetttaagt 2340
taaatatgta gatcaaggat gtccaatctt ttggcttccc tgagccacac tggaagaaga 2400
attotettoo geegeacata aaatatgeta acaetgatga tagetgat
<210> 61
```

<211> 2255 <212> DNA <213> Homo sapiens

<220>

<221> misc-feature <223> Incyte ID No.: 320087CB1

PCT/US00/02237

```
<400> 61
ttccqqttct gtaccccat cctttctctc gcccttcta cccgcagetc ctqqcqctcq 60
geggggetaa etgeageggg gagatetegg eegecaaget eegecteeeg eecegggetg 120
tgccccgggg ctcgcctgag gccgaccacc cgcaacccac ctctagcggc tttgctcgag 180
geometric trecacees oggommetre cagtaggets geometre actoroges 240
cccgcgtcaa ctgcaagggg cccgcccata gccagttccg gggcggttgc tcacatcgac 300
eggaacteee egeceettee eqeggeeeet ggggeegtag gaggeegeag egaggaggta 360
gaggggggg gggtcgcact agggtgtccc tagagaacga ggactctgaa ggcgggacat 420
ttgggcgacc cccgggcggg gccagccatt aaacagtccc acttctgtgc cagacactga 480
actgggetet tgacgggeat catetettaa teeteagaac ateccaggga getecacagg 540
atccccatat cotgggccat gagtgagttg aaagactgcc cottgcagtt ccacgacttc 600
aagtetgtgg atcacetgaa ggtetgteee egetacaegg cagtgetgge aegetetgag 660
gatgatggea teggeatega ggagetggae accetgeage tggagetgga gaccetgetg 720
tettetgeca geeggegeet gegtgtgett gaggeegaaa cecagateet cacegaetgg 780
caqqataaga aaqqtqacaq acqattcctq aagctgggtc gagaccatga acttggagct 840
cceccaaac atqqqaaqec caaqaaqcaq aaactggaag ggaaggcagg acatgggccg 900
ggccctggcc caggacggcc caaatccaaa aaccttcagc ccaagatcca ggaatatgaa 960
ttcactgatg accetatega cgtgecaegg atceccaaaa atgatgecce caacaggite 1020
tgggetteag tggageeeta etgtgetgae atcaccageg aggaggteeg cacacttgag 1080
gagttactga agcccccaga agatgagget gagcattaca agatcccacc cctggggaag 1140
cactactece agegetggge ceaggaggae etgetggagg ageagaagga tggggeegg 1200
gcagcggctg tggctgacaa gaagaaaggc ctcatggggc cactgaccga actggacact 1260
aaagatgtgg atgccctgct gaagaagtct gaggcccagc atgaacagcc ggaagatgga 1320
tgcccctttg gtgccctgac gcagcgcctc ctgcaggccc tggtggagga aaatattatt 1380
teccetatgg aggattetee tatteetgae atgtetggga aagaateagg ggetgaeggg 1440
gcaagcacct cccctcgcaa tcagaacaag cccttcagtg tgccgcatac taagtccctg 1500
gagageegea teaaggagga getaattgee eagggeettt tggagtetga ggacegeee 1560
gcagaggact ccgaggatga ggtccttgct gagcttcgca aacggcaggc tgagctgaag 1620
quantitaging cocacaaceq caccaaqaaq cacqaccinc thaggering aaaggaggag 1680
gtgagccggc aggagctgag gcagcgggtg cgcatggctg acaacgaggt catggacgcc 1740
tttcqcaaqa tcatqqctqc ccqqcaqaaq aaqcggactc ccaccaagaa agaaaaggac 1800
caggectgga agactetgaa ggagegtgag agcateetga agetgetgga tgggtageec 1860
teaccectge eteaggetga ttatetggee taggggaggg gaagggagge ceactteett 1920
ctttgggcac aggaaacatt ggcctgtggc tgtccctcaa atggcggcag tctctagagg 1980
geographics treecting arctititing characterist acaaccagga cacaggaage 2040
cctgctgggc tagcctgagg cctagtctct gcttggtccc cgagatgggg ttggagggga 2100
cttegtttet gggtetteet etteceetet ttaccatece ccacteceta atcccetace 2160
cctgtctccc cttcaaggac ttctcccttg tggttttgta aagtgcaaac ttaagaataa 2220
agtgactgct gtggtttttc aaaaaaaaa aaaaa
                                                                    2255
<210> 62
<211> 2982
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incyte ID No.: 491271CB1
gegtggggat gtttacegee gtttateegg gatagagact ceategtget gacageatee 60
ttttattcac cgcctccgaa tttgcaaaga ggaggaagga gggacttstt ggsttctccc 120
agcatagece cagttatgee ateteagaae tatgacette eecagaagaa geaggagaaa 180
atgaccaagt ttcaggagge tgtgacatte aaggatgtgg ctgtggtett ctccagggag 240
gaactgcgac tgctcgatct tacccagagg aagctgtacc gagatgtcat ggtggagaac 300
tteaagaace tggttgeagt ggggeatett ceetteeaac cagatatggt ateceaattg 360
gaagcagaag aaaagctttg gatgatggaa acagaaaccc aaagaagcag caagcatcaa 420
aataagatgg aaacactcca aaaatttgca ttaaaatacc tttcaaatca agagetgtcc 480
tgctggcaaa tctggaaaca ggttgcaagt gaattaacca ggtgtcttca ggggaagagt 540
teccagetat tacaaggega etetatteag getteetgaaa algagaacaa talaatgaac 600
cctaaaggag atagccctat ttatattgaa aatcaagagt ttccattttg gagaacccag 660
```

PCT/US00/02237

```
cattettqcq qqaatacata tetqaqtqaq teacagatte agagtagaqq taaqeaaatt 720
gatgtgaaaa ataacctgca aatacgtgaa gacttcgtga agaaatcacc atttcatgag 780
catattaaaa ctgacacaga accaaaaccc tgcaaaggta atgaatatgg caaaatcatt 840
agtgatggct ccaaccagaa attaccctta ggagagaaac cccatccatg tggtgagtgt 900
ggaaggggct teagttatag eccaaggett ecectteate egaatgttea cacaggagaa 960
aaatgettea gteaaagete acatetgega acteateaga gaatteacee aggagagaaa 1020
ctcaatagat gtcatgaatc tggtgattgc ttcaataaga gctcttttca ttcttatcaa 1080
totaatoata caggagagaa gtottataga tgogacagtt goggcaaggg attoagtago 1140
agcacgggtc ttatcattca ttacagaact catactggag agaaacccta taaatgcgag 1200
gaatgtggta aatgctttag tcaaagttca aattttcagt gccatcagag agtccacact 1260
gaagaaaaac catacaaatg cgaagagtgt ggtaagggct tcggttggag tgttaatctc 1320
cgtgttcacc agagggtcca caggggtgag aagccctata aatgtgagga atgtggtaag 1380
ggetteacte aggetgeaca ttttcacate catcagagag tecacactgg agagaaacce 1440
tacaagtgtg atgtgtgtgg taagggcttc agccacaatt caccattaat atgccatcgg 1500
agagtecaca caggagagaa qecatacaag tqtgaggegt gtgggaaagg ctttacccgt 1560
aatacaqatc tqcatattca tttcaqaqtt cacacqqqag aqaaacccta taaatgtaag 1620
gagtgtggta agggettcag tcaggettca aatettcaag tccatcagaa tgtccacact 1680
ggggagaaac gattcaagtg tgaaacgtgt gggaagggct tcagtcagtc ctcaaagctt 1740
gacttcagtt atagttcaaa tottaaacta caccaagtaa ttcacactgg agaaaaacca 1860
tataaatgtg aggaatgtgg gaagggcttc agttggagat caaatcttca tgcacatcaa 1920
agagtteact caggagaaaa accetataaa tgtgagcagt gtgataagag etteagteag 1980
gccatagatt ttcgggtaca tcagagagtc catactggag agaagccata caaatgtggt 2040
gtotgtggta agggottcag toagtoctot ggtottcaat cocatcagag agtocacacg 2100
ggggaaaagc catacaaatg tgatgtgtgt ggaaagggct ttagatacag ttcgcagttt 2160
atataccatc agagaggesa cactggagaa aaaccttaca aatgtgaaga gtgtgggaaa 2220
ggctttggta ggagettgaa tettegeeat cateagaggg tecacaeggg agagaaacce 2280
catatatgtg aggagtgtgg taaggcottc agtotoccot caaatottog agtocacotg 2340
qqtqttcaca ccaqqqaaaa actctttaaa tqtqaaqaqt qtqqtaaaqq cttcaqtcaq 2400
agtgcacqtc ttqaaqccca tcagagaqtc cacactggag aaaaaccata caaatgtgac 2460
atatgtgata aggaetteeg teacegttea egtettacat ateateagaa agtecataet 2520
ggtaaaaagc tttagaaatg agaaatgtgt taccaacttt tgtctgaatg cacatcttca 2580
agttttttggc tagtccatgc tggtggtaaa ccctgtaaaa ctactgagag tggaaggggg 2640
tttgttcaca cttggaatct ttctaacaaa tccatcaaga tgataacaca gaaccatgaa 2700
caggaataga actcgtattt aggggagaaa tagggctggt ggctctcttg gtaagatcta 2760
gttaatataa atgatcacct ttcattgtga atatatgcct gaagataatg tgtggaagga 2820
tatttgccat atgctaactg gtttttggc cagggagagt tttgggttat tatccctttt 2880
ctttaatttt cattttatac ttacagtgat cattattttc ataaaagctg taaagctatg 2940
aaaaatgaat aaaattacta aaaattttct gtaaaaaaaa aa
                                                                  2982
<210> 63
<211> 1185
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incvte ID No.: 585172CB1
ggagcgtcca gagtcctggc cctgagcggg aatcgcagtg gccgaggctg agcggcaggt 60
agaagggggg tctccggggc ttcacaggga acacaggggc ttcggcccaa ccacaagtac 120
geagttgeac atgeectact tttgeecatt ctttgeaaat cccctaagga acgaacgege 180
ctogogtgog ggoettteta atettegett gtecetteae ttecacaget ggaggtegaa 240
tteaccaegt caegtagaga aaaggggogg tgtetegggt eteecegtgt ggeatcaagg 300
cgggctccct ataggagctg gtgtggacgt ccgggcgtgg ggttagggcg gatgcgggga 360
tegeggggeg ggtgttgaca ttgtgetete accaggegga tegeceegae ceteactect 420
ggegtetgag tetetggegt agceatgetg agtgggegge tggteetggg tetggtetee 480
atggctggcc gcgtttgttt gtgccagggc agcgcgggat ccggggccat cggtccggtg 540
gaggeegeea ttegeacgaa gttggaggag geeetgagee eegaggtget agagettege 600
```

aacqaqaqeq qtqqccacqc qqtcccgcct qqcaqtqaqa ctcacttccg cgtggctgtg 660

建氯化物甲基苯二酚 建筑 "不明,我就是我们的我们的人,就是不知识的人。"

PCT/US00/02237

```
grangetete grittegaggg actgaggece ctacaacgae accggetggt ecacgeageg 720
ctggccgagg agctgggagg tccggtccat gcgctggcca tccaggcacg gacccccgcc 780
cagtqqaqaq agaactetca getgqacaet agesecceat geetgggtgg gaacaagaaa 840
actctaqqaa cccctqaac cccaaqaqaq qqaqqaccaq qatccqaatg ggctgggtga 900
geacgaatta ccgaggeett ccctttgata cagtecagga tttgtaaggg atgaagaccc 960
ctgggcccca ttctgttggg gtccatacat actctccgaa gatagcaact tgcttcaggt 1020
caaaqtqaac ccqaqaaaaq aqaaqaatca ctcactactq ctcttqccct qqactattca 1080
ggaagggeag cccggatgtt ccatgttaaa tcgtgacaga attgcaccag acctgatgag 1140
ttggaaacaa tcctatacat taaaagaaat racactaaaa aaaaa
<210> 64
<211> 1191
<212> DNA
<213> Homo sapiens
-220×
<221> misc-feature
<223> Incyte ID No.: 615200CB1
<400> 64
tggacggacg cgtgggcttg atttctgatt tatgactgct ttttgttgta ccccaatagt 60
cqtctaaqaa aqqtqattat tttgaqaggc ctggggagac acacatgctc attctcqagq 120
gtggcggtgg tgcagagggc agagccatge tegttettge tateetgaga ttggttgeta 180
totqtttcct ttqctgctgt gtttttttct gtcagtatta aaggtggaag aaggtccata 240
tettttetg tgggtgette aagtgttgtt ggaagtggag geageagtga caaggggaag 300
ctttccctgc aggatgtagc tgagctgatt cgggccagag cctgccagag ggtggtggtc 360
atggtggggg ccggcatcag cacacccagt ggcattccag acttcagatc tccggggagt 420
quectigaca quaaceteca quagtacgat etcoogtace cogaggocat ttttqaacte 480
ccattettet tteacaacce caageesttt tteactttgg ccaaggaget gtaccetgga 540
aactacaage ccaacatcac tcactacttt ctccggctgc ttcatgacaa ggggctgctt 600
ctgcqqctct acacqcaqaa catcqatggg cttgagagag tgtcgggcat ccctgcctca 660
aagctggttg aagctcatgg aacctttgcc totgccacct gcacagtctg ccaaagaccc 720
tteccagggg aggacatteg ggetgaegtg atggeagaea gggttecceq etgeeeggte 780
tgcaccggcg ttgtgaagcc cgacattgtg ttctttgggg agccgetgcc ccagaggttc 840
ttgctgcatg tggttgattt ccccatggca gatctgctgc tcatccttgg gacctccctg 900 gaggtggagc cttttgccag cttgaccgag gccgtgcgga ctcagttccc cgactgctca 960
tcaaccqqqa cttqqtqqqq cccttqqctt ggcatcctcg cagcagggac gtggcccagc 1020
tgggggacgt ggttcacggc gtggaaagcc tagtggagct tctgggctgg acagaagaga 1080
tgcgggacet tgtgcagcgg gaaactggga agcttgatgg accagacaaa taggatgatg 1140
getgececca cacaataaat ggtaacatag gagacateca cateccaatt c
<210> 65
<211> 2596
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 997067CB1
<400> 65
coggtgtgtg ggtctgtgac agggtccaac agggcctggt ccgtgtccgg tcccccaaat 60
ctgtcgtccc tgccccagg cattggcatc aacaaaagtc agaattcccg ggaacttgaa 120
cagaggetge taaatteeca gtaattgete etttggeett etagggaetg aetteaaaga 180
aggaaggaaa gaatcaggca gtgcttcctc attctcttt aaaacccgct tcccgctgag 240
totgoaccca ggagaccaga gagcacettg coettocatg gaaactcagg etgatetegt 300
atctcaggaa cctcaggccc tgcttgacag tgctcttcct tcaaaaagttc ctgccttttc 360
cgacaaggac agcotggggg atgagatgtt ggcggctgcg ctcctaaagg ccaagtccca 420
ggagetggta acctttgagg atgtagetgt gtacttcate eggaaggagt ggaagegttt 480
ggaacetget cagagggace tetatagaga tgtgatgetg gagaattacg ggaatgtgtt 540
```

62/91

中国美国工作。1962年118日 - 中国工作中国工作中国工作中国工作。

STAN BUSY OF BURNESSES WITH

PCT/US00/02237

```
ctcactggat cgtgagacca ggactgaaaa tgatcaagaa atttctgaag acacaagatc 600
acatggggtc ctactgggaa gatttcaaaa ggatatttct cagggtctca agtttaaaqa 660
agectatgaa cgagaagtca gtetgaaaag geegetgggg aacteeeetg gagaaagaet 720
gaacaggaaa atgccagatt ttggtcaagt gacagttgag gagaagctaa cccccagggg 780
agagagaage gagaaatata atgattttgg gaacagette actgtgaatt ccaacettat 840
ctcacatcag agactccccg tgggagacag accccataag tgtgatgaat gtagcaagag 900
ctttaatcga acttcagacc ttattcaaca tcagagaatc cacactgggg aaaagcccta 960
tgaatgtaat gagtgtggga aggeetteag ceagagetea caeettatte ageateagag 1020
aatccacact ggggaaaaac cttatgaatg tagtgattgt gggaaaacct tcagctgtag 1080
ctetgecete attetgeate ggaggateca caegggggag aaaccetatg aatgtaatga 1140
gtgtgggaag accttcaget ggagetecae ceteacecae cateagagaa tecacaetgg 1200
tgagaaaccc tacgcctgca atgaatgtgg gaaggccttc agcaggagct caacccttat 1260
tcaccatcag agaatccaca ctggagaaaa accctatgaa tgtaatgaat gtgggaaagc 1320
cttcagccag agetcacace tetatcagca ccagagaate cacactggag agaageceta 1380
cgaatgtatg gaatgtggag gaaagtttac ctacagttca ggccttattc agcatcaaag 1440
aatccacacc ggggagaacc cctatgaatg tagtgagtgt gggaaagcct tcaggtacag 1500
ctcggctctt gttcgccatc agagaattca cactggagag aagcctttga atgggatcgg 1560
catgagcaaa agctccctca gagttacgac cgagttaaat atcagagagt ccacgtgaaa 1620
gagccacaca cccattttcc tcactttccc tgagtctcaa gagctcttgc cttaccctat 1680
aaatotcaac agottaggat gtgtcccttt caactcagac ttttcatttt agagaatggg 1740
geagatgggg caaategttg aatttteeca gaaateacac cageettaga aagegteaag 1800
gcaagtggat ggcgtgctgg gaatagaaag cagctctggg accagttacc ccatttagga 1860
aaggagtttg cactaaactg tttttctcac agcagaggaa cctttccaag gtggggatgg 1920
aagcacagte gggacagaat tegatggatt cetttagttg gagteegegt agteagcaca 1980
gacagcagtg gaaaggacgc totgggtoot gttatttgct agggagggta aagggagact 2040
atttcaaagc tactgttcct agtccagctt taagtttcgg taagaaacat gctgttttgt 2100
gagattccag aatcatcatc tgtgatgatg gtgtccttta gggctcttgg agcagccaga 2220
ccatqtttcc aaqaqaaacc tggtgatatt gccagcagac cccctgccat cccccccagt 2280
tgtcctgggg ctgaatgggc aaatctgtcc aaacagctag taaccggctg tgagggagag 2340
ggtcagaage acttagegtt ggcctctgat tgctgtcctc tcttgtcctc ttcccactcc 2400
aatgatgaaa atgattttet etaaatgeet gggtaaggat gettteaagg ageteaettg 2460
geotgettge cotgecotot cacetetgae acceagecee aggagecaga ecacteetge 2520
ctccacctct gactcttcag cagctgaaga ttaatgcaga gaaagagcaa agcccaaaag 2580
                                                                 2596
ggagaaaaaa aaaaaa
<210> 66
<211> 1574
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incvte ID No.: 1443262CB1
cagecegtga gaegeceget getggaegeg ggtageegte tgaggtgeeg gagetgeggg
aggatggage cgctgaaggt ggaaaagttc gcaaccgcca acaggggaaa cgggctgcgc 120
gccgtgaccc cgctgcgccc cggagagcta ctcttccgct cggatccctt ggcgtacacg 180
gtgtgcaagg ggagtcgtgg cgtcgtctgc gaccgctgcc ttctcgggaa ggaaaagctg 240
atgegatget etcagtgeeg egtegecaaa tactgtagtg etaagtgtea qaaaaaaget 300
tggccagacc acaageggga atgcaaatgc cttaaaagct gcaaacccag atatcctcca 360
gactccgttc gacttcttgg cagagttgtc ttcaaactta tggatggagc accttcagaa 420
tcagagaagc tttactcatt ttatgatctg gagtcaaata ttaacaaact gactgaagat 480
aagaaagagg gcctcaggca actcgtaatg acatttcaac atttcatgag agaagaaata 540
caggatgeet eteagetgee acetgeettt gacetttttg aageetttge aaaagtgate 600
tgcaactett teaceatetg taatgeggag atgeaggaag ttggtgttgg cetatatece 660
agtatetett tgetcaatca cagetgtgac cecaactgtt cgattgtgtt caatgggeec 720
cacctettac tgcgagcagt ccgagacatc gaggtgggag aggagetcac catctgctac 780
```

ctggatatge tgatgaceag tgaggagege eggaageage tgagggacea gtactgettt 840 gaatgtgact gttteegttg ecaaaeceag gacaaggatg etgatatget aactggtgat 900

LANGE OF FULL AND A

PCT/US00/02237

```
gagcaagtat ggaaggaagt tcaagaatcc ctgaaaaaaa ttgaagaact gaaggcacac 960
tggaagtggg agcaggttet ggccatgtgc caggcgatca taagcagcaa ttctgaacgg 1020
cttcccgata tcaacatcta ccagctgaag gtgctcgact gcgccatgga tgcctgcatc 1080
aacctcggcc tgttggagga agccttgttc tatggtactc ggaccatgga gccatacagg 1140
atttttttcc caggaageca tecegteaga ggggttcaag tgatgaaagt tggcaaactg 1200
cagctacate aaggcatgtt teeccaagca atgaagaate tgagaetgge ttttgatatt 1260
atgagagtga cacatggcag agaacacage ctgattgaag atttgattet acttttagaa 1320
gaatgcgacg ccaacatcag agcatcctaa gggaacgcag tcagagggaa atacggcgtg 1380
tgtctttgtt gaatgootta ttgaggtcac acactctatg ctttgttagc tgtgtgaacc 1440
tetectatty gaaattetgt teegtgtttg tgtaggtaaa taaaggeaga catggtttge 1500
aaaccacaag aatcattagt tgtagagaag cacgattata ataaattcaa aacatttggt 1560
tgaaaaaaa aaaa
<210> 67
<211> 2197
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 1521648CB1
<400> 67
cccqacqaac gcgaggaggg cgaggcggga ggtgcaggag ggaccctcgc catgggtcca 60
cgggcctaga gtggcggaag ataccggcct ggtgccaaac tggctactgc tgcttcctgt 120
ggcctccatg gctgaggact ggctggactg cccggccctg ggccctggct ggaagcgccg 180
cgaagtettt cgcaagtcag gggccacctg tggacgetca gacacctatt accagagece 240
cacaggagac aggatccgaa gcaaagttga gctgactcga tacctgggcc ctgcgtgtga 300
totoaccoto ticgactica aacaaggoat ottgigotat coagoccoca aggoccatec 360
cgtggcggtt gccagcaaga agcgaaagaa gccttcaagg ccagccaaga ctcggaaacg 420
tcaggttgga ccccagagtg gtgaggtcag gaaggaggcc ccgagggatg agaccaaggc 480
tgacactgac acagececag etteatteee tgeteetggg tgetgtgaga actgtggaat 540
cagettetea ggggatggea eccaaaggea geggeteaaa aegttgtgea aagaetgteg 600
agcacagaga attgccttca accgggaaca gagaatgttt aagcgtgtgg gctgtgggga 660
gtgtgcagcc tgccaggtaa cagaagactg tggggcctgc tccacctgcc tcctqcaqct 720
gccccatgat gtggcatcgg ggctgttctg caagtgtgaa cggagacgct gcctccggat 780
tgtggaaagg agccgagggt gtggagtatg ccggggctgt cagacccaag aggattgtgg 840
ccattgeece atetgeette geecteeceg eeetggtete aggegeeagt ggaaatgtgt 900
ccagegacgt tgcctacggg gtaaacatgc ccgccgcaag ggaggctgtg actccaagat 960
ggctgccagg cggcgcccg gagcccagcc actgcctcca ccaccccat cacagtcccc 1020
agageceaca gageegeace ceagageeet ggeeceeteg ceacetgeeg agtteateta 1080
ttactgtgta gacgaggacg agctacagcc ctacacgaac cgccggcaga accgcaagtg 1140
cggggcctgt gcagcctgcc tacggcggat ggactgtggc cgctgcgact tctgctgcga 1200
caageccaaa ttegggggca gcaaccagaa gegecagaag tgtegttgge gceaatgeet 1260
geagtttgee atgaagegge tgetgeecag tgtetggtea gagtetgagg atggggeagg 1320
ategececea cettacegte gtegaaagag geceagetet gecegaegge accatettgg 1880 cectacettg aageceacet tggetacaeg cacageceaa ceagaecata eccaggetee 1440
aacgaagcag gaagcaggtg gtggctttgt gctgcccccg cctggcactg accttgtgtt 1500
tttacgggaa ggcgcaagca gtcctgtgca ggtgccgggc cctgttgcag cttccacaga 1560
agccctgttg caggtgaagc aagagaaggc ggatacccag gacgagtgga caccaggcac 1620
agetgteetg actteteegg tattggtgee tggetgeeet ageaaggeag tagacceagg 1680
cctgccttct gtgaagcaag agccacctga cccagaggag gacaaggagg agaacaagga 1740
tgattetgee tecaaattgg ceccagagga agaggeagga ggggetggea caccegtgat 1800
cacggagatt ttcagcctgg gtggaacccg cttccgagat acagcagtct ggttgccaag 1860
gtccaaagac cttaaaaaac ctggagctag aaagcagtag actggaggct tctacagact 1920
gaccgaggac acagtggagc ccacgagcac gagctggaac ccacgaggat ggcctggaac 2040
ccatgleagt eteteaceae etecagette gatgatgtgg gtgteetgea gaagaagetg 2100
gtgcccttcc tcacagagtt aaatatgcat ctggcccagg aattagagaa gctgaaagga 2160
tgatectggg gaaggtggaa cagetgeagg cetgget
```

PCT/US00/02237

```
<210> 68
<211> 2081
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incvte ID No.: 1685494CB1
<400> 68
gcatacgcag teegtgeeta ageggeacga ecategagee aaggteetee agegtteeta 60
gageggagaa gaaagegete egaagageta gagetgacae teggegatga getaagaege 120
tgtttcagag cgtttgggtc ctctgaggcc ccttgaccag gagtgtctct gaagatacag 180
tecaaagaag gttetecaaa acaaggagag cagtetgaag etggggatgg caacagcatt 240
ggtgagtgcc cattccctgg ctcccctgag tctgaagaag gaggggcttc gggtagtgag 300
ggaggatcac tactctactt gggaacaggg attcaagctg caaggaaaca gtaaaggcct 360
tggacaggag ccattgtgca aacaattcag gcagttgcgt tatgaagaga ccacaggacc 420
tcgagaagca ctaagtcggc tccgggagct ctgtcaacag tggctacagc ccgagaccca 480
taccaaggag cagatcctgg agctgctggt gctggagcag tttctgatca tcctqcctaa 540
ggagetecag geeegggtge aggageatea eccagagage agggaggaeg tggttgttgt 600
totggaggat ttgcagctgg atottggaga aacaggacaa caggtggacc cagaccagcc 660
aaagaaacaa aaaatacttg tggaggagat ggcccctctg aaaggagtac aggaacagca 720
ggttcggcat gagtgtgaag ttacaaagcc tgagaaagag aagggtgagg agacaaggat 780
aatatetgaa cecatggagg eteataatga gggetetaae ttggaaagge ateaggecaa 900
geccaaagag aagattgagt ataaatgete agaaegtgag cagagattea tecageaett 960
ggacctgatt gaacatgcga gtacacacac gggaaagaaa ctctgcgagt ctgatgtgtg 1020
tcagagttcc agtcttacag gacataagaa agtcctctct agagagaaag gtcatcagtg 1080
tcatgagtgt gggaaagcct ttcagaggag ttcacacctc gtcagacatc agaaaatcca 1140
tettggtgag aageettate agtgcaatga gtgtggcaaa gtetttagee agaatgcagg 1200
ccttttggaa catctcagaa ttcatactgg agagaaacct tatctatgta tccattgtgg 1260
aaaaaatttt aggcgcagct ctcaccttaa tcgacatcag agaattcaca gtcaggagga 1320
gccctgtgag tgcaaggagt gtggaaaaac ctttagtcag gccttactcc tcacccacca 1380
tcagagaatc catagtcact ccaaaagcca tcaatgtaac gagtgtggaa aagctttcag 1440
tttgacctca gaccttattc gacaccacag aattcatact ggagaaaaac ctttcaagtg 1500
taacatatgc cagaaagcct teegactaaa etcacacett geteagcatg taagaateca 1560
caatgaagaa aaaccctatc agtgtagtga atgtggagaa gccttcaggc aaaggtcagg 1620
tettttteaa cateagagat atcaccacaa agacaaactg gettgatgag gtgttetete 1680
cttgtagaac atcagagaag gcacattgac tagcaaacag cactttagga aaagtcaccg 1740
tageccactg tggcatcaga aaattettgg gggetgagtt ggaggeteee tgeetetatt 1800
ctctctcctt tgctttcctt gaagtcagct ttggaccaca ataatttcac tgtagatgat 1860
atgctaggat caaagttaaa cagcattett caetgcagga catetcagag catgtaacat 1920
aactgcatga ttatatactc taagcaatag agagetteat gactgagtaa gagttttgaa 1980
gtcagcagtg aatcaagtge ccacagattt gcaggettaa gcagaacaag ggaagattga 2040
tatttttgga tatgetatag cagetttete ctatgaaata a 2081
<210> 69
<211> 2785
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incvte ID No.: 1730829CB1
<400> 69
gaccgtgcgg cgcccagcgg agtaggggct gcgcttgggg tttgctgaag ctqqctgcct 60
ctcccactcc ccttttgggt gcaaagcgcc gctagcggga agacgggggc cgggcgggga 120
caggggcacc tgcgtactgg actgagagcc tgcgcccagc ttacatcgac cccacccggc 180
cccggcccga cccgacgcga cccgatccga tccgatccca ttccatccgt tcctcgtctc 240
ctcccggtct gacccgttgc ccggccgtgg ttcgccacac caggcatcca aagctgaggt 300
```

65/91

the war to an in the first that I have the

PCT/US00/02237

```
egetectacg geotgggete geottegett tagagatgtt tggeetette ceteccaaac 360
arcteatett caaaacetqq actettqqae tggcacetgg ccacetttee etctaccaag 420
actocactte egtettacce acttetteet cagattettg gtaccecetg ggttggagac 480
tgctcatttt ccttccaaat taatcccaga ccccctaaaa tattgacaac cttgacaacc 540
ceccaacega ggagecagae tttgttttgg actaacttcc atagecetat catggaggea 600
gtgtacctgg tagtgaatgg gttgggcctg gtgctggacg tgctgacctt ggtgttggac 660
ctcaacttcc tgctggtgtc ctccctcctg gcttccctgg cctggctcct ggccttcgtc 720
tacaacctgc cgcacacggt actgactagt cttctgcact tgggccgcgg agtcttgctt 780
teattgetgg cettgatega ageegtggte eggtteacat gtgggggett geaggeettg 840
tgtactetgc tgtatagetg ctgetetggc ctagagagec taaageteet ggggcacetg 900
geeteteatg gggeactgeg gageagggag atactgeace ggggegteet caatgtggte 960
tecagtggee atgetttget gegeeaggee tgtgacatet gtgeeattge catgageetg 1020
gtggcttatg tgatcaacag cctggtcaac atctgcctca tcggcactca gaacctcttt 1080
tecetggtge tggeeetgtg ggatgeagtg accgggeete tgtggaggat gacagaegta 1140
gtggetgeet teetageeca catttecage agtgetgtgg ccatggecat ceteetttgg 1200
acaccetgee aactageeet ggagetgetg geeteagetg coegeeteet ggeeagettt 1260
gtgcttgtca atctcactgg cttggtgttg ctagcttgtg tgctggcagt gacggtgact 1320
gtgttgcatc cggacttcac cctgaggctg gctacccagg cactcagcca gctccatgcc 1380
eggecatect accacegtet tegagaggat gteatgegge tetetegeet ageaetggge 1440
tcagaggeet ggegeegagt etggageege agtetgeage tggegagttg gecaaacegg 1500
ggaggggcac ctggagctcc ccagggtgac cctatgaggg tattctcagt taggacccgg 1560
agacaggaca ctcttcctga agcggggcgc agatcagagg cagaagagga ggaggccagg 1620
accatcagag tgacacctgt caggggccga gagaggctca atgaggagga gcctccaggt 1680
gggcaagace cttggaaatt getgaaggag caagaggage ggaagaagtg tgtcatetge 1740
caggaccaga gcaagacagt gttgctcctg ccctgccggc atctgtgcct gtgccaggcc 1800
tgcactgaaa tcctgatgcg ccaccccgtc taccaccgca attgcccgct ctgccgccgg 1860
ggcatcctge agaccctcaa tgtctacctc tgaagcctcc ttccctgcct gcccacccct 1920
ccatgeteca egeaggeact caegetagga cageattaac aceteatete egggteetgg 1980
totgaatoco etectacece tgtggccate etgecataca tecaggacat tgagttggaa 2040
gactatgate tgggtggggg caggataaca tggettetet ttacccagtg ggtecetteg 2100
atgctgaggg tggtgagtat gtcactatgc aagggeeetg agactatttg etgtgggete 2160
tectecagee tgeccaggge ccaeccagat geetetgggg ttacceetgt etgectetgg 2220
tttttctgtt ggagatctat aggtcctttt cctgcctcct tcacatttcc tccccagett 2280
ttygggcac acacatcag tgtcatttgg gtgttttgg aactcagggg cttcggatg 2340 atcttaaacc ttytgttta gccagagcc cttgtgccctg gtaggcgtt gggttagta 2400 cttcgggtg ccctcagagc cactctcg ttggtgttgtg 2400 acctcagagtg
ctgatccaaa agccagtete aggagtttac ccctgggatg qgggatgcat ctgcacctga 2520
etttggggee acgtgeeetg tggcacceca getcactggg agtetcagga gggataaccg 2580
gatttetget ettteecetg teacteceae atcacacaga aaaatggeat teetetetgt 2640
ctctccctgg catggagagg gcagactgtg cacatttcac tagggttcaa atacagaagg 2700
cccagggccc aggggttgca gcttcgtgag gggtctctgg cccagtttcc aatgaataaa 2760
                                                                     2785
gttctcttga cagctaaaaa aaaaa
```

```
<210> 70
```

<400> 70

tcccgtccta	tgacgtcagc	cgtaaggcgc	tgctgtcgta	aaaggacgtc	cggtccgtct	60
cctagtgtcc	ggaatcggct	gtcagcctcc	ctggctgtta	gtaccttctt	tcccggagtc	120
ctagtccaca	agttggattt	actgetgteg	cgggtgggcc	tcacgccatt	ccctgtccct	180
caaccccta	agtgagtccg	ateteccaac	gaaagtgagc	gaggtttgcc	cggagcgcgc	240
acgagggaa	aatgcctaaa	aaaaagactg	gtgcgaggaa	gaaggetgag	aaccgccgag	300
aacgtgaaaa	acaactaaga	gcatcaagaa	gcactataga	tttagctaaa	catccatgta	360
					ttttgctact	

<211> 1231 <212> DNA

<213> Homo sapiens

<220>

<221> misc-feature

<223> Incyte ID No.: 1864641CB1

PCT/US00/02237

```
tttgtaatte tgtacagaag ttaccaattt gtgcacagtg tgggaaaaca aagtgcatga 480
tgaagtotto agactgtgto ataaagcatg otggtgtata cagtactggo ottgcaatgg 540
tgggtgcaat atgtgacttc tgtgaagctt gggtttgcca tggtaggaaa tgtctcagta 600
cacatgettg tgeetgeect ettacegatg etgagtgtgt tgaatgtgaa egaggegtgt 660
gggaccatgg aggcagaata ttcagttgtt ctttttgcca taactttctc tgtgaagatg 720
atcaatttga gcatcaagcc agctgccagg ttttagaggc agaaacattt aaatgtgttt 780
catgcaatcg gettggtcag cactcatgte teegttgtaa ggettgttte tgtgatgate 840
atacaaggag caaagtgttt aagcaagaaa aaggaaaaca gcctccttgt cctaaatgtg 900
ggcatgaaac teaggagact aaggacetta gcatgteaac acgetecetg aaatttggca 960
ggcagactgg aggtgaagag ggagatggag cttctgggta tgatqcttat tggaagaacc 1020
tttcatctga taagtatggt gataccaget accaegatga ggaggaggat gagtatgaag 1080
cagaggatga tgaagaggaa gaagatgaag gcagaaagga ttcagatact gagtcatcag 1140
atttqtttac taatttqaat ttaggaagga cctatgctag tggctatgct cactatgagg 1200
aacaagagaa ctaggggage tgctctggtg g
                                                                     1231
<210> 71
<211> 700
<212> DNA
<213> Homo sapiens
<220> -
<221> misc-feature
<223> Incvte ID No.: 2444604CB1
<400> 71
gccccaggtg acacaatggc cgcagtccat ggcggctggc ttcttccagc ccttcatgtc 60
accordette ccaqqqqqcc cccqqcccac cctqcqqatq ccqaqtcaqc ctcccqcatq 120
cotcoctgge teccageece tectecetgg egecatggag coetececae gageecaggg 180
geateegage atgggegge caatgeagag ggtgaegget cetegtggea tggceagegt 240
ggggccccag agctatggag gtggcatgeg acccccaccc aactccctcg ccggcccagg 300
cotgectgcc atgaacatgg geccaggagt tegtggeceg tgggccagec ccagtggaaa 360
ctcgatccc tactcctcct catcccccgg cagctacacc ggacccccag gaggaggtgg 420 gcccctgga acacccatca tgcctagccc tggagattcc accaactcca gcgaaaacat 480
gtacactate atgaacccca tegggeaggg egeeggeagg getaatttee egeteggeee 540
tggcccggag ggcccatggc cgccatgagc gcgatggagc ctcaccacqt gaacqqatcc 600
ctgggctcgg gcgacatgga cgggttgccq aagaqttccc ccggcqccqt gqccgqcctq 660
agcaacgccc cggggcaccc cgcgggacga cggcgagatg
<210> 72
<211> 2332
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2445008CB1
aggeggaegg ggaaegagge egteggeeat tttgtgtetg etteetgtgg gaegtggtgg 60
tagecettgg gttgggaaag tgagggattt ttggeetegt tteteetget tetttetee 120
teeettttae tittgeeggta gaacacagtt atgggtegea agaaqaaqaa qeaqetgaaq 180
ccgtggtgct ggtattgtaa tagagatttt gatgatgaga agateettat tcagcaccaa 240
aaagcaaagc attttaaatg ccatatatgt cacaagaaat tgtatacagg acctggctta 300
getatteatt geatgeaggt acataaagaa acaatagatg cegtaccaaa tgcaatacet 360
ggaagaacag acatagagtt ggaaatatat ggtatggaag gtattccaga aaaagacatg 420
gatgaaagac gacgacttct tgaacagaaa acacaagaaa gtcaaaaaaa gaagcaacaa 480
gatgattetg atgaatatga tgatgaegae tetgeageet caaetteatt teageeaeag 540
cetgttcaac ctcagcaagg ttatattcct ccaatggcac agccaggact gccaccagta 600
ccaggageac caggaatgee tecaggeata cetecattaa tgecaggtgt tectectetg 660
```

67/91

atgccaggaa tgccaccagt tatgccaggc atgccacctg gattgcatca tcagagaaaa 720

· California Swa Adam no sa

PCT/US00/02237

```
tacacccaqt cattttgcqq tqaaaacata atqatqccaa tqqqtqqaat qatqccacct 780
 ggaccaggaa taccacctct gatgeetgga atgecaccag gtatgeeccc acctgtteca 840
 cgtcctggaa ttcctccaat gactcaagca caggctgttt cagcgccagg tattcttaat 900
 agaccacctg caccaacage aactgtacct geoccacage etecagttac taagectett 960
 ttccccagtg ctggacagat ggggacacct gtcacaagct caagtacagc ttcatccaat 1020
gctgtccaag gacctgttgg tacagatttc aaacccttaa atagtacccc tgcaacaact 1140
acagaaccc caaagcctac attecetget tatacacagt ctacagette aacaactagt 1200
acaacaaata gtactgcagc taaaccagcg gcttcaataa caagtaagcc tgctacactt 1260
acaacaacta gtgcaaccag taagttgate catecagatg aggatatate cetggaagag 1320
agaagggcac agttacctaa gtatcaacgt aatcttcctc ggccaggaca ggcccccatc 1380
ggtaatccac cagttggacc aattggaggt atgatgccac cacagccagg catcccacag 1440
caacaaggaa tqaqacccc aatgccact catggtcagt atggtqqtca tcatcaaggc 1500
atgccaggat accttcctgg tgctatgccc ccgtatgggc agggaccqcc aatggtgccc 1560
cettaccagg gtggggeetee tegaceteeg atgggaatga gaceteetgt aatgtegeaa 1620
ggtggccgtt actgatetta etteatecag tetaataggt ttggagatta aacettttet 1680
caacttgtgc tgtttatata gccaagettc cgtcaataag gcttcattgt gactttaaca 1740
aacattatct tcccacatac caggaactat tggacattta ttttacatgg gaaaaattat 1800
ttggaataat aaagcaggaa cttttcctga agttgcaatt tatactgtat ggcttctttt 1860
tcatgtttca tctaggtttt tagaagtgaa gtatagtaaa tttggttcgt taaattgtga 1920
aggogotgga attacatgaa cataccacco tagtaaaggo aagttotgta agottacatt 1980
getatttgta aagtttgeet teacagcatt teagatgetg ttggaettea tgteeccaae 2040
ctagettggt gagggetgta actgtttcca agtacttgta cattggaagt etgaatgtgt 2100
aacaatattt aatgtattta gagtteetea tgttgeaggg tttaagaaat etgaceeace 2160
aaggtcatgt gacttttctg tactgttaaa cttcattgta ataaaatgag agaaaaattt 2220
atgeettttt atteataace cagetgtgga ceaetgeetg aaaggtttgt acagatgeat 2280
<210> 73
<211> 1936
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incyte ID No.: 2572462CB1
ttggttctat attactgttt tatagctgag gagaaagact cacatcattt actaagatca 60
tatagetage tagtaaatgt ttgagtgaaa atacaaacaa aggttttetg actttaagag 120
cttgagtttt ttccactata ccatattgca tctqttgtaa ttqttaacta atqtqcattt 180
taaaattete attigtetta tgtactgage eettatacea gtgetaattt atgtgaetee 240
tttctcctgc agctaagaga aaaatacctt tttaattcat ttatagtacc cagtttttaa 300
agaagattta tittgtaaaa tittgettat ggtacatgte atettageet gtaaataaat 360
taaagcatta atttttatcc ctccctggtc ttttcctcct tctgacttta tacgtctttc 420
tagagagett atettetata ataacaatte tttgttttaa agtgagaaag ateagtetaa 480
agaaaaggag aagaaagtga aaaaaacaat teetteetgg getaceettt etgecageca 540
gctagccagg gcccagaaac aaacaccgat ggcttcttcc ccacgtccca agatggatgc 600
aatcttaact gaggccatta aggcatgett ccagaagagt ggtgcatcag tggttgetat 660
tegaaaatac atcatecata agtateette tetggagetg gagagaaggg gttateteet 720
taaacaagca ctgaaaagag aattaaatag aggagtcatc aaacaggtta aaggaaaagg 780
tgcttctgga agttttgttg tggttcagaa atcaagaaaa acacctcaga aatccagaaa 840
cagaaagaat aggagetetg cagtggatee agaaccacaa gtaaaattgg aggatgteet 900
cccactggcc tttactcgcc tttgtgaacc taaagaagct tcctacagtc tcatcaggaa 960
atatgtgtct cagtattatc ctaagcttag agtggacatc aggcctcagc tgttgaagaa 1020
cgctctgcag agagcagtag agaggggcca gttagaacag ataactggca aaggtgcttc 1080
ggggacattc cagetgaaga aatcagggga gaaacccctg cttggtggaa gcctgatgga 1140
atatgcaatc ttgtctgcca ttgctgccat gaatgagccg aagacctgct ctaccactgc 1200
```

68/91

tetgaagaag tatgteetag agaateacee aggaaceaat tetaaetate aaatgeattt 1260 getgaaaaaa accetgeaga aatgegaaaa gaatgggtgg atggaacaga tetetggggaa 132 agggtteagt ggeacettee agetetgitt tecetattat eccageceag gagttetgit 1380

```
tccgaagaaa gagccagatg attctagaga tgaggatgaa gatgaagatg agtcatcaga 1440
agaagactet gaggatgaag agecgecace taagagaagg ttgcagaaga aaaccecage 1500
caagteecca gggaaggeeg catetgtgaa geagagaggg tecaaacetg cacetaaagt 1560
ctcagctgcc cagcqqqqqa aaqctaqqcc cttqcctaaq aaaqcacctc ctaaqqccaa 1620
aacgcctgcc aagaagacca gaccctcatc cacagtcatc aagaaaccta gtggtggctc 1680
ctcaaagaag cctgcaacca gtgcaagaaa ggaagtaaaa ttgccgggca agggcaaatc 1740
caccatgaag aagtetttea gagtgaaaaa gtaaatttta taggaaaaaa gggtateatg 1800
atgaaattca aaatcttatt ttctaaggtc agtgtgcatt totttagttt tgatgctttt 1860
caaattacat tattttcctc ccctatgaac attgtgggga gggactctaa ataaaccagt 1920
ttaggcaaaa aaaaaa
<210> 74
<211> 1667
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incvte ID No.: 2572892CB1
<400> 74
egegaategg egaccecagt geetegacca ctatgeegeg etettteete gteaggaage 60
cctccgaccc caatcggaag cctaactaca gcgagctgca ggactctaat ccagagttta 120
cettecagea gecetacgae caggeecace tgetggeage cateceacet ceggagatee 180
teaacceae egectegetg ceaatgetea tetgggaete tgteetggeg ceceaaggee 240
agecaattge etgggeetee etteggetee aggagagtee cagggtggea gagetgacet 300
ccetgtcaga cgaggacagt gggaaagget cccageceee cageceacee teaceggete 360
cttegteett etectetaet teageetett cettggagge egaggeetat getgeettee 420
caggettggg ccaagtgee aageagetgg cccagetete tgaggecaag gatetecagg 480
ctegaaagge ctteaactge aaatactgea acaaggaata ceteageetg ggtgeeetea 540
agatgcacat ccgaagccac acgetgccct gcgtctgcgg aacctgcggg aaggcettet 600
ctaggccctg gctgctacaa ggccatgtcc ggacccacac tggcgagaag cccttctcct 660
gtccccactg cagccgtgcc ttcgctgacc gctccaacct gcgggcccac ctccagaccc 720
actcagatgt caagaagtac cagtgccagg cgtgtgctcg gaccttetec cgaatgteee 780 tgctccacaa gcaccaagag teeggetget caggatgtee eegetgacee tegaggetee 840
cicticetet ceatacetge ceetgeetga cageetteee cagetecage aggaaggace 900
ccacatcett eteactgeca tggaatteee teetgagtge eccaettetg gecacateag 960
ccccacagga ctttgatgaa gaccattttc tggttctgtg tcctctgcct gggctctgga 1020
agaggeette eegtggeeat ttetgtggag ggagggeage tggeeceeag eeetggggga 1080
tteetgaget ggeetgtetg egtgggtttt tgtateeaga getgtttgga taeagetget 1140
ttgagetaca ggacaaagge tgacagacte actgggaage teccaceca etcaggggac 1200
cocactocco toacacaca cococcacaa ggaaccotca qqccaccotc cacqaqqtqt 1260
gactaactat gcaataatcc accccaggt gcagccccag ggcctgcgga ggcggtggca 1320
gactagagtc tgagatgccc cgagcccagg cagctatttc agcctcctgt ttggtggggt 1380
ggcacctgtt tcccgggcaa tttaacaatg tctgaaaagg gactgtgagt aatggctgtc 1440
acttgteggg ggeceaagtg gggtgetetg gtetgaeega tgtgteteee agaactatte 1500
tgggggcccg acaggtgggc ctgggaggaa gatgtttaca tttttaaagg tacactggta 1560
tttatatttc aaacattttg tatcaaggaa acgttttgta tagttatatg tacagtttat 1620
tgatattcaa taaagcagtt aatttatata ttaaaaaaaa aaaaaaa
<210> 75
<211> 759
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incyte ID No.: 2785674CB1
<400> 75
```

renderation of the state of the

PCT/US00/02237

```
tgggctcgcc tccctgggac taggtttcag cggccgctgc gatgaccaaa ataaaggcag
atcccgacgg gcccgagget caggcggagg cgtgttccgg ggagcgcacc taccaggage 120
tgetggteaa ccagaaccec atcgegcage ccctggcttc tcgccgcctc acgeggaage 180
tctacaaatg catcaagaaa gcggtgaagc agaagcagat tcggcgcggg gtgaaagagg 240
ttcagaaatt tgtcaacaaa ggagaaaaag ggatcatggt tttggcagga gacacactgc 300
ccattgaggt atactgccat ctcccagtca tgtgtgagga ccgaaatttg ccctatgtct 360
atatececte taagaeggae etgggtgeag eegeaggete caagegeeee acetgtgtga 420
taatggtcaa gccccatgag gagtaccagg aggcttacga tgagtgcctg gaggaggtgc 480
agtccctgcc cctaccccta tgaggggctc cggtagcacc tgggcacctg ccgctggaag 540
ctattggget ggeageagga egactggetg tecteetgee cacceacact gaeggeatet 600
teccagttee ccaaggeacg cettetteee aggeagetet aacageeett teatgaaggt 660
aatgctagtc ttctgtccat cagtgccatt tcctgtagaa ctaaaggctg ttccaagaat 720
gtggggtggg gaaagtaaat gctaagacta aaaaaaaa
<210> 76
<211> 1421
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2797479CB1
<400> 76
ccccagteac tgttcctgcc cgttgcctgt cagcctcact gcctttaatc cgcttccagg
getcattcae etgaatatat tggttgeeat gtegagatat ggtgaggata etgegeaagg 120
catcocggat ocgaacotto toaagtaget gttgatagtg otggagetco acggtgacat 180
gagctaacag gcgctgatca tcatgggtga tgacatatcg acgcacatag cccccaaaga 240
acttagacac aaacatccca getetgttga sgaagttgee caggttgtta ageageteag 300
aattattett cagcagcagg teegteeagg agaaagcaet gteetggeee teaggeegaa 360
tgtacagcag atagaagcgc cagatgtcag cagggatccc cgtgtcctgg gctttgattt 420
tectggttea geacgaatte atgaaggaae teacacteta gagaaaeeet atgaatgtaa 480
gcaatgtggg aaattgttat ctcatcgctc aagctttcga agacacatga tggcacacac 540
tggagatggc cctcataaat gcacagtatg tgggaaagcc tttgactctc ctagtgtatt 600
tcaaagacat gaaaggactc acactggaga gaaaccctat gaatgcaagc aatgtgggaa 660
agcetteegt aetteeagtt ceettegaaa acatgaaaca acacacatg gagageaacc 720
ctataaatgt aaatgtggaa aagcttttag tgatttattt tcctttcaaa gtcatgaaac 780
aacacacagt gaagaggagc cttatgaatg taaggagtgt gggaaagcat ttagttcttt 840
taaatacttt tgtcgccatg aacggactca cagtgaagaa aaatcttatg agtgtcaaat 900
ttgtggcaaa ctttcagtcg tttcagttac ttaaaaactc atgaaaggac tcacacggca 960
gagaagccat atgaatgtaa gcaatgcagg aaagcattct tttggccctc tttccttcta 1020
agacatgaaa ggactcacac tggagaaaga ccctatgaat gtaaacactg tggtaaagcc 1080
ttcagtcgtt ccagtttctg tcgagaacat gaaagaactc acgctggaga gaagccctat 1140
gaatgtaagg aatgtgggaa agcettcagt tetetcagtt cetttaatag acataaaagg 1200
acacactgga aggatattet ataagtgtat ggaatgtggg aaagcattea ttggttttat 1260
cacattcaga tacttgaaag aaataaatcc tgtgaatgta aacgtggtaa agccttaaga 1320
agtttccagg ctgggcgcag cggctcacac ctgtaatccc agcactttga gaggccgagg 1380
agggcatate acgaggccag gagategaga ceageetggg g
 <210> 77
 <211> 2386
 <212> DNA
 <213> Homo sapiens
 <220>
 <221> misc-feature
 <223> Incyte ID No.: 2960640CB1
 <400> 77
 gacttttaag ggatcacaga gctcacacca aagaccaggg gaacagtcag aagcctggct 60
```

PCT/US00/02237

```
tgeteeteag geteeeagga acetgeetea aaacacaggt etecacqace aggagacaqq 120
tgctgtggte tggacagetg ggteccaggg accagecatg cgtgacaaca gagetgtate 180
cetetgteag caagaatgga tgtgeeeagg ceetgeacaa agggeeetet acaggggtge 240
cacccagagg aaggacagte acgteteget ggcaacaggt gtgccetggg getatgaaga 300
gaccaagacg ctcctggcta ttcttagtag ttctcaattt tatggaaaac tccagacctg 360
tcagcagaac agccagatct acagggccat ggcggaagga ctctgggagc agggttttct 420
geggaccea gaacagtgte geaccaagtt caaaagecta cagttgagtt accgcaaagt 480
gaggagagge egtgtgeetg ageettgtat ettttatgag gaaatgaatg etettteagg 540
ctcctgggcc tctgcacctc ctatggcaag cgatgctgtt cctggccaag aaggaagtga 600
tattgaggct ggagagctga atcaccagaa tggggaaccc acggaggtag aagatggcac 660
tgtggatggt gcagacaggg atgaaaagga cttcaggaat cctggccagg aagtcaggaa 720
actagacctg ccagtgctgt tcccaaacag acttggtttt gagttcaaga acgagattaa 780
aaaagaaaat ctaaaatqqq atqattcaga qqaaqtaqaa ataaacaagg ctttacagag 840
aaagtccaga ggagtttatt ggcactctga gctacaaaaa ggcttggaga gtgagccaac 900
atcaagaagg caatgtagaa attctccagg ggagagtgag gagaaaaccc catcccagga 960
gaagatgagt caccagagtt tttgtgccag ggacaaagcc tgtacacata tcctctgtgg 1020
gaaaaactgc tetcagagtg tgcactetec ccacaageca gegetcaaac tggaaaaagt 1080
atctcaatgt cetgaatgtg ggaaaacett tageegaagt tettatettg tteggeatea 1140
aaqaatccac acaggcgaga agcetcacaa gtgcagtgag tgcgggaagg getttagtga 1200
gegetecaae eteaetgeee acetaegaae teacacaggg gagaggeeet ateagtgtgg 1260
qcaatqtqqq aaaaqcttca accagaqttc cagcctcatt gtccaccaga ggacccatac 1320
cggggaaaag cottaccagt gcattgtotg tggaaagaga ttcaacaaca gttcccagtt 1380
cagtgctcac cggcgcatcc acactgggga gagcccatac aagtgtgcag tgtgtgggaa 1440
aatetteaac aatageteec aetteagtge ceaecgaaaa acceaeactg gtgaaaagec 1500
ttacaggtgt teteactgtg agagaggett cactaagaac tetgecetca eccgtcatca 1560
gacagração atgaaaggag tactorcato acaggaagga agagatgcgt tatgagtgtg 1620
toggtaaact gtcagattaa gttcctcagg tcagcatgta tgagctttct tctgctgtgg 1680
agagatetag ceagteeetg aetttgeaac agacetaetg aetaetggat ettaagaeee 1740
atgictagga ccaggagtca gcataaggac gctgacctct cctggctgtg cctgtgactc 1800
cagagtecta tettactgtg acttaaagtt tgatggagag aaagetgtag gateeataaa 1860
ttctaccagg aatccagggt cttcctgttc ctagcactga gaatgggcac ccagtggtcc 1920
aaqaacactt tetgggetaa catagteete acacaggget gagaaagaaa gtgteteett 1980
tcctggaaag cacatgtaaa agttaagggc ctgaatctct tctaaaccaa taattgacct 2040
ctaggcaccc accgtaatac tgctaccttc agagcagaag acactgctct ttgtaaccca 2100
gccaccacca aaaagcaaac agaaagaagg aaggatggtt aagccattgg ataatactga 2160
atctqtttcc ctaagtgact taaccttaga gcaagcacaa catccaggtt aattaattgt 2220
aaqatttete eteteattat tgeecteate atagtteetg attgtetett aaagtaagtg 2280
gtttatagac attactattt ctgataataa tttacaaact actataaaca aatttataaa 2340
cattactaat ttctgatgaa aataaagttg tttctccctc caccac
                                                                  2386
<210> 78
<211> 1432
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incyte ID No.: 3454051CB1
aqtatgaqct qqacctqtcc gcgttqccag caacctgttt tcttcgctqa gaaggtgagc 60
tecetgggea agaactggea ecgettetge etgaaatgtg agegetgeea eageateetg 120
teccetggeg ggeatgeaga geacaatggg aggeeatact geeacaagee atgetatggg 180
getetetttg gacceagggg ceeteeceat atgaagacat teaetgggga gacetegetg 240
tgccctggct gtggggagcc cgtctatttt gctgagaagg tgatgtcatt aggcagaaat 300
tggcaccgac cgtgtctgag gtgccagcgt tgccacaaga ccctgactgc tgggagtcat 360
getgageatg atggagteec ctactgeeac gteceetget aeggetacet gtttggeece 420
aaaggtgtga acattggcga tgtgggctgc tacatctatg acccagtgaa gataaaattc 480
agatgagacg etcacaaaaa aggtcaccct aactcaggec teccatcatg ecectcatgg 540
```

tocaatggaa gotacaaaaa totocagtoo catgggggtt gggggaaggtg ggatottggg 600 ggcotgggoo taggotocat ggtaggooag agagtotaga otttototgo caatttttt 660

<213> Homo sapiens

```
cettteccat ttetatetgg ttagggaaca gecegttttg aagggtatee teeteetgge 720
catcacaacc Cottteccca geacattetg gagetteaag gtacteataa aacttgtgtt 780
 tattgaattt cagoctetgt ggettettea ataaaatgtt ggeteecatg cetteaacte 840
ttetttggge atgagaceag tgggttggag gatggggagt gtgggggttg ggatgacatg 900 cattgccetg cagggtgcet cggaggtage agggccagce atgagaacaa aaagetetgt 960
tetttttgte cettgggeet ggeattggea gteetageae cacacagtgg acagcatgee 1020
caccagecce attggtacca gaagteteat atgetagtee tttetttage accateteta 1080
gaagaagcag aagcacctta ttcagtaact catttgagca tggcaacaga tcctatggta 1140
gggcctccca agaggcttca ttatccattt gggagatgaa cagactgagg cccagagagg 1200
gaaagccaca tgcccaaggt cacacagcaa gttaatggtg aaggttttat cagagcccag 1260
ggcagactca gtggcttcct atccagggct cttctcacag ctcgtcacca ctgccccaac 1320
ccaaggggca cctttattta cagaatctcc ccaaccgtga gacgggtgcc agcagaccac 1380
tgcattctgg gaagtcaact gttcctagaa gcaaataatc aaggatggat ga
<210> 79
<211> 1816
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incvte ID No.: 3510640CB1
<400> 79
cactgatgca ggaactgtat agcacaccag cotocaggct ggactcottc gtggctcagt 60
ggctgcagcc ccaccgggag tggaaggaag aggtgctaga cgctgtgcgg accgtggagg 120
agtttctgag gcaggagcat ttccagggga agcgtgggct ggaccaggat gtgcgggtgc 180
tgaaggtagt caaggtqqqc teetteqqqa atqqcacqqt teteaggaqc accaqaqaqq 240
tggagctggt ggcgtttctq agctgtttcc acagcttcca ggaggcagcc aagcatcaca 300
aagatgttct gaggctgata tggaaaacca tgtggcaaag ccaggacctg ctggacctcg 360
ggctcgagga cctgaggatg gagcagagag tccccgatgc tctcgtcttc accatccaga 420
ccagggggac tgcggagccc atcacggtca ccattgtgcc tgcctacaga gccctggggc 480
cttotottec caacteccag ccaccectg aggtetatgt gageetgate aaggeetgeg 540
gtggtcctgg aaatttctgc ccattcttca gcgagctgca gagaaatttc gtgaaacatc 600
ggccaactaa gctgaagagc ctcctgcgcc tggtgaaaca ctggtaccag cagtatgtga 660
aagccaggte ceecagagee aatetgeece etetetatge tettgaactt etaaccatet 720
atgcctggga aatgggtact gaagaagacg agaatttcat gttggacgaa ggcttcacca 780
ctgtgatgga cctgctcctg gagtatgaag tcatctgtat ctactggacc aagtactaca 840
cactccacaa tgcaatcatt qaggattgtg tcagaaaaca gctcaaaaaa qagaggccca 900
tcatcctgga tccggccqac cccacctca acqtqqcaqa aqqqtacaqa tqggacatcq 960
ttgctcagag ggcctcccag tgcctgaaac aggactgttg ctatgacaac agggagaacc 1020
ccatctccag ctggaacgtg aagagggcac gagacatcca cttgacagtg gagcagaggg 1080
gttacccaga tttcaacctc atcgtgaacc cttatgagcc cataaggaag gttaaagaga 1140
aaateeggag gaceagggge tactetggee tgeagegtet gteetteeag gtteetggea 1200
gtgagaggca gcttctcagc agcaggtgct ccttagccaa atatgggatc ttctcccaca 1260
ctcacateta tetgetggag accateceet eegagateea ggtettegtg aagaateetg 1320
atggtgggag ctacgcctat gccatcaacc ccaacagctt catcctgggt ctgaaqcagc 1380
agattgaaga ccagcagggg cttcctaaaa agcagcagca gctggaattc caaggccaag 1440
tectgeagga etggttgggt etggggatet atggcateca agacagtgae acteteatec 1500
totogaagaa gaaaggagag gototgttto cagocagtta gttttototg ggagacttot 1560
ctgtacattt ctgccatgta ctccagaact catcctgtca atcactctgt cccattgtct 1620
actgggaagg teccaggtet teaccagttt tacaatgagt tateccagge cagacgtggt 1680
ageteacace tgtaateeca gaactttggg aggeegaggt gggaggageg ettgageega 1740
ggagttcaag accagectgg gtatcatagg gagaccccqt Ctctacaaaa taaaaaaata 1800
attcactggg aaaaaa
                                                                   1816
<210> 80
<211> 1556
<212> DNA
```

```
-2205
<221> misc-feature
<223> Incyte ID No.: 3815083CB1
<400× 80
ctcaggteeg gagegeggte gggacacage geetetagga gaaageetgg aaggegetee 60
gggggtatcc agagctetta gcgggccggc agcatgtgcg gggccccagt aaatggaaat 120
gttttctaac atataaaaac ctacagaaga agaaaataat tttctggatc aaattagaag 180
tetgtattat attgatgtet ecagatteaa atatattaga aageageegt ggagacaace 240
atottoattt tgggagaaat aactaaagoo ogootoaago attagaacta cagacaaaco 300
ctgatgegac ctctccagat tgtcccaaqt cgattgattt cccagctata ttgtggcctg 360
aageeteeag egteeacaeg aaaceagatt tgeetgaaaa tggeteggee aagtteaagt 420
atggcagati ttcgaaagti ttttgcaaaa gcaaagcaca tagtcatcat ctcaggaget 480
ggtgttagtg cagaaagtgg tgttccgacc ttcagaggag ctggaggtta ttggagaaaa 540
tggcaagece aggacetgge gactecett geetttgee acaaccegte coggetgtgg 600
gagttctacc actaccggcg ggaggtcatg qqqaqcaagg aqcccaacgc cgggcaccgc 660
qecatageeg agtgtgagac ceggetggge aagcagggee ggegagtegt ggtcatcace 720
cagaacateg atgagetgea cegeaagget ggeaceaaga acettetgga gatecatggt 780
agettattta aaactegatg tacetettgt qqagttgtgg ctgagaatta caagagteca 840
atttgtccag ctttatcagg aaaaggtgct ccagaacctg gaactcaaga tgccagcatc 900
ccagttgaga aacttecceg gtgtgaagag gcaggetgeg ggggettget gegaceteae 960
gtcgtgtggt ttggagaaaa cctggatcct gccattctgg aggaggttga cagagagctc 1020
geocactgtg atttatgtet agtggtggge actteetetg tggtgtacce ageagecatg 1080
tttgcccccc aggtggctgc caggggcgtg ccagtggctg aatttaacac qqaqaccacc 1140
ccagetacga acagattcag gtttcatttc cagggaccct gtggaacgac tcttcctgaa 1200
gcccttgcct gtcatgaaaa tgaaactgtt tcttaagtgt cctggggaag aaagaaatta 1260
cagtatatet aagaactagg ceacacgcag aggagaaatg gtettatggg tggtgagetg 1320
agtactgaac aatctaaaaa tagcctctga ttccctcgct ggaatccaac ctgttgataa 1380
gtgatggggg tttagaagta gcaaagagca cccacattca aaagtcacag aactggaaag 1440
ttaatteata ttatttggtt tgaactgaaa cgtgaggtat etttgatgtg tatggttggt 1500
tattgggagg gamaaatttt gtaaattaga ttgtctaaaa aaaataaaaa aaaaaa
<210> 81
<211> 1951
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 3988457CB1
ctccgattat gggataggag aagtgcccgt ggagcccctg gatgtcccct taccctccac 60
gatcaggeca getteeceeg tggcegggte tecaaageag eeggtgegtg getactaceg 120
tggcgctgtc ggtggcacgt ttgaccgcct gcacaacgcc cacaaggtgt tgctcagtgt 180
cgcgtgcatc ctggcccagg agcagcttgt ggtgggagta gcagacaaag atctgttgaa 240
gagcaagttg ctccctgagc tgctccaacc ttatacagaa cgtgtggaac atctgagtga 300
attectggtg gacatcaage cetecttgac ttttgatgtc atcecectge tggaccecta 360
tgggcccgct ggctctgacc cctccctgga gttcctggtg gtcagcgagg agacctatcg 420
toggggggatg gccatcaacc gcttccgcct tgagaatgac ctggaggaac ttgctttgta 480
ccagatccag etgetgaagg acctcagaca tacagagaat gaagaggaca aaqtcagete $40
ctccagcttc cgccagcgaa tgttggggaa cctgcttcgg cctccatatg aaaggccaga 600
geteceeaca tgtetetatg taattggget gaetggeate agtggetetg ggaagagete 660
aatageteag egaetgaagg geetggggge gtttgteatt gaeagtgace aectgggtea 720
tegggeetat geeceaggtg geeetgeeta ceageetgtg gtggaggeet ttggaacaga 780
tattetecat aaagatggca teateaacag gaaggtesta ggeageeggg tgtttgggaa 840
taagaagcag ctgaagatac tcacggacat tatgtggcca attatcgcaa agctggcccg 900
agaggagatg gategggetg tggetgaggg aaagegtgtg tgtgtgattg atgeegetgt 960
gttgcttgaa gccggctggc agaacctggt ccatgaggtg tggactgctg tcatcccaga 1020
gactgagget gtaagaegea tigiggagag ggatggeete agigaageeg eggeteaaag 1080
coggetgeag agceagatga gegggeagea gettgtggaa cagageeaeg tggtgeteag 1140
```

the beautiful and the same of the same of

```
cagecettgt gggageegea tateacceaa egecaggtgg agaaageetg ggceetettg 1200
cagaagegea ttcccaagac tcatcaggec ctcgactgaa aagttetcag tggggccaga 1260
ctggctcctg gagctgacaa gcgaccccqt ggtgaggaga aatgggggcc ttgatgctca 1320
ccctggttca ggcccagagg tccaagctat actgtgcagg acatggccag gcctggtgga 1380
cacaggaage ctacccaaca coctootatt tooccaacac toaggatoto ottcatgggg 1440
gagcagtece etecceaete tigeceatgg gigactetta eccaeagetg actagggeea 1500
gegeaaatac tggaacctgt aacagaatta aaggtgaatg ttetgaaaaa aaaatagaat 1560
tttggacatc tacaactaac tcgatttaca cttacgaaca taatggactc ttaaaaaatg 1620
gaaagggata acagggaccc cccgggttct gcatccttcc tcccggggat ttttttccgg 1680
coggeteett gegggtgaac tgatttteet tacactgege etatttaaac gttggggtaa 1740
ccagggtegg acctitteec tiggaactit tittaccegt cacgatitec actgeaacti 1800
agacteggga aacttagggt ggaateeetg gggggeetaa ggegggaact eeeteeettt 1860
atotgotttg ggtcaaagag cocgtttotc catotggcaa ctotgtccat ccaaggggtt 1920
ttcgggttct cgggccaagg cccggggtgg g
<210> 82
<211> 1313
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incyte ID No.: 131890CB1
egtegggagg geetaagtee gtgtgeggtg eeetteggee ggeetgagee ecagagteag 60
ctecettte tegeceageg ecceaggee geteeegggg etcaeggaat agtaaagaaa 120
cacatcataa aacctcccag gacataaagg tgagcacaga ccctgtttgg atcaagtcag 180
tteetggage etgaatgatg aetgetgaat caegggaage caegggtetg teeccacagg 240
ctgcacagga gaaggatggt atcgttatag tgaaggtgga agaggaagat gaggaagacc 300
acatgtyggg geaggattee accetacagg acacgcetee tecagaccca gagatattee 360
gecaacgett caggegette tgttaccaga acaettttgg geceegagag getetcagte 420
ggetgaagga actitgteat cagtggetge ggecagaaat aaacaccaag gaacagatee 480
tggagettet ggtgetagag eagtitetti ceateetgee caaggagete caggietgge 540
tgcaggaata ccgccccgat agtggagagg aggccgtgac ccttctagaa gacttggagc 600
ttgatttatc aggacaacag gtcccaggtc aagttcatgg acctgagatg ctcgcaaggg 660
qqatqqtqcc totqqateca qttcaqqaqt cotcqaqctt tgaccttcat cacgaggcca 720
cccagtccca ettcaaacat tegtetegga aacceegeet ettacagtca egaggtaaga 780
agcaaggttt catttagggg aagggaaatg attcaggacg agagtetttg tgctgctgag 840
tgcctgtgat gaagaagcat gttagtcctg ggcaacgtag cgagaccca tctctacaaa 900
aaatagaaaa attagccagg tatagtggcg cacacctgtg attccagcta cgcaggaggc 960
tgaggtggga ggattgcttg agcccaggag gttgaggctg cagtgagctg taatcatgcc 1020
actactecaa eetgggeaac acagcaagga eeetgtetea aaagetaett acagaaaaga 1080
attaggeteg geaeggtage teacacetgt aateceagea etttgggagg etgaggeggg 1140
cagatcactt gaggtcagga gtttgagacc agcctggcca acatggtgaa accttgtctc 1200
tactaaaaat atgaaaatta gccaggcatg gtggcacatt cctgtaatcc cagctactcg 1260
qqaqqctqaq qcaqqaqaat cacttqaacc caqqaqqtqq aqqttqcaqt aag
<210> 83
<211> 1197
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 238642CB1
<400> 83
eggetegage gtgaacegaa gteettatat teeegggett ettteteete tgggtaceag
ctecttactg ccctgcagac aagcgtgccg tgcgtgcttg tggccaaggg aaggaagagc 120
tggttgatcc acagatagct cettectecc egeceettee titttgtttg gaggteccag 180
```

PCT/US00/02237

```
gatctgtgtt cacagacatc tgggggaaga aaaggagcag gaaactaccc cgcacagagt 240
taagcaggaa acaacaacaa catcatgcaa aaaccctgca aagaaaacga aggaaagcca 300
aagtgcagcg tgccaaagag ggaggaaaaa cgcccgtatg gagaatttga acgccagcaa 360
acagaaggga attttagaca gaggetgett cagteteteg aagaatttaa agaggacata 420
ttggaagaga taaggggtot gagaaagaaa tttagggoto tgcattotaa ccataggoat 540
tetegggace gteettatee catttaatta atttetetga caatteaatt attttetgtt 600
attaatgttg ceactgettt etgtttgtet geactttett gataaatatt tgetategtt 660
ttacteeagt eattegatgt tgetgagatt tacatatgae tettgteaae ateteatett 720
ttgacccaat cttattcatt taataagagg tctcattcat ttgcatggaa aaatgctcat 780
tgtatattgc aaagtgaaaa taacgagttg caaaacagtg tatacatata tgtgtgtata 840
tatgtacact ttatttgtac atttctatgt gacataatgc aaaggaaagt gtctgatttt 900 attatacacc aaaggttaac agtgaatctc tgtgtgatct ctttttttt ctttttgcct 960
atctgcatct teteacttge caaaaaatga atatatgttt atgtgtgtat attacttgtg 1020
tcacaaaaaa ccctaaagta gacagtaaaa gaacttgtca atcgcctttg gaaggcaatg 1080
asacacttaa taaactctca ataacagaag cgtaaaaatg aaatgtaaac ctccaattac 1140
ctctqqatct cttaqccaqa qtaataaact qqtaattatt acaqataaaa aaaaaaa
<210> 84
<211> 2170
<212> DNA
<213> Homo sapiens
-220×
<221> misc-feature
<223> Incyte ID No.: 669862CB1
<400> 84
ageqtqeqeq ataqqacaqq geettaaatt catqqttatq tqttttgqttt tcattctaag 60
ataaagccga accactgaca aaattatata gagagcaatg tgataatgtt aggaatagag 120
atattcaqqa catactcctq tqcattctqa tccactcaqt ctqaqtqqat atctaaaaqa 180
tagtgtatgt aaaatgetee tatggtaatt etgatttgea actaaatatg agaactagtg 240
actitigtett tgeetaetea taaatettae ateeteetgt geactitieat aattitietgt 300
gecettettq cettttatat attgettgge agtactttag aattttetaa actetttgat 360
tactaacaga tacacttttt agttgatcac tatttttaat atagatgtcc tctccctacc 420
cgctattatt ggaaaatagc atttgtttgt tttttcattt tcttccagac tttaatttca 480
caactgaaag caacaagtta tottoagaaa aaagaaatta tgaagtaaat gogtaccato 540
aggagacatq gaaaagaaat aaaaccttca accttatgag gtttattttc agaactgacc 600
cacaqtacac aattqaattt qggagacaac aqagacctaa agtqqgatqt tttagtcaaa 660
tgatattcaa aaaacataaa teeetteete tacataagag aaataacaca agagagaaat 720
catatgagtg taaggaatat aagaaggget ttagaaaata tttgcacctt actgaacatc 780
tgagagacca tactggtgtg ataccctatg aatgtaatga atgtggaaaa gcatttgtag 840
ttttccagca ttttattaga catcgaaaaa tccacactga tttgaaaccc tatgaatgca 900
atggatgtga gaaggeettt aggttttatt cacagettat teagcateag ataatteata 960
ctggtatgaa accctatgaa tgtaagcaat gcgggaaggc ttttagacgt cattctcacc 1020
ttacagaaca tcagaaaatt catgttggct tgaaaccctt tgaatgtaag gaatgtgggg 1080
aaacgtttag attatatcga catatgtgtc tgcatcagaa aattcatcat ggtgtgaaac 1140
cctacaaatg taaagaatgt ggaaaggett ttggtcateg ttcaagtett taccaacata 1200
agaaaattca ttetggtgag aaaccatata aatgtgaaca atgtgaaaag gcetttgtte 1260
geagetatet aettgttgaa catcaaagaa gteataetgg tgagaaacet catgaatgea 1320
tggaatgtgg aaaggetitt agtaaggget caageettet taaacataag agaatteata 1380
gtagtgagaa actetatgat tgtaaggatt gtggaaagge ettttgtaga ggeteteaac 1440
ttacacagca tcagagaatt catactggtg agaagccaca tgaatgtaaa gaatgtggga 1500
agacttttaa getteattea tatettatte aacateagat aatteataet gatttgaage 1560
catatgaatg taagcaatgt gggaaagcct tcagtcgtgt tggagacctt aagacacatc 1620
aatcaattca tgctggggag aaaccctatg aatgtaagga atgtggaaaa acctttagac 1680
ttaattotca actaatttat catcagacaa ttcatactgg tttgaaaccc tatgtatgta 1740
aagaatgtaa gaaggeettt egttetatet caggtettte teaacataag agaatteata 1800
ctggtgaaaa accctatgaa tgtaaagaat gtgataaggc ctttaatcgc agtgatcgac 1860
ttactcaaca tgagacaatt catactggtg tgaaaccaca gaaatgcaaa gaatgtggta 1920
aggeetttag teattgetat caacttagte aacateaaag attteaceat ggtgagagae 1980
```

tal and a few to day while characterists have resonanced the contribution of the same time.

WO 00/44900 PCT/US00/02237

```
tottaatgta atgagaggga aagcotttag coatggoaca tttttactgt tgtcactatt 2040
atgatgctat agtgaagatt aaactagtta atatagaata taaatacttg gaaaggcatc 2100
tggcacatcg tatttgctta ctaaatacat tatttttatg ataattgtta gaattactaa 2160
                                                                    2170
gaataaatga
<210> 85
<211> 1904
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incyte ID No.: 1003663CB1
<400> 85
cageteteag gteegacace egetggaage eggegegge geaggegege aegeaaagge
ggccgggagt aaggcggact gaaggaggag cttgatggaa gcgtgcgaga aggggcgtaa 120
ctgatttgga aaccagagga aaggcgctgt tttcaccgaa ttagaatcgc gggaaaatag 180
agaagagttt gtttgaaggt ctcgcqagat cgaqaccgga agtccttcat ctcaagcatc 240
caatgetgaa ageggeetga ttttetetae eggaageeet ttteeagagg etgggaacae 300
ggcccaccta gcaggaagte ccacctectt gageteegee accetteeeg aagttittet 360
greacctgtg traggered teceetrice gegittiate eccgraceag aaaaggatae 420
atttagtgee teccacecag etccactaaa egggttggat ateteattet ttgagttggt 480
gtteetteee eggegeeeee atgtagetgg gaagtgggae etgggggtgg ttggaeeeet 540 gggateetaa aggaggggea gggagggege agaaeteege ttetgeteet tgetaceagg 600
acgogogoc tectcageet etttectece getgecatge accetgeage ettecegett 660
cetgtggttg tggccgctgt getgtgggga gcggcccga cccgggggget cattcgagcg 720
accteggace acaatgecag catggacttt geagacette eagetetgtt tggggetace 780
ttgagecagg agggeteca ggggtteett gtggaggete acccagacaa tgeetgeage 840
cecattgee caccacece ageceeggte aatgggteag tetttattge getgettega 900
agattegact geaactttga ceteaaggte etaaatgeee agaaggetgg atatggtgee 960
getgtagtac acaatgtgaa ttccaatgaa ettetgaaca tggtgtggaa tagtgaggaa 1020
atccagcage agatetggat eccgtetgta tttattgggg agagaagete egagtacetg 1080
egtgeeetet tigtetaega gaagggget egggtgette tggtteeaga caatacette 1140
cccttgggct attacctcat ccctttcaca gggattgtgg gactgctggt tttggccatg 1200
ggagcagtaa tgatagctcg ttgtatccag caccggaaac ggctccagcg gaatcgactt 1260
accaaagagc aactgaaaca gattcctaca catgactatc agaagggaga ccagtatgat 1320
gtctgtgcca tttgcctgga tgaatatgag gatggggaca agctgcgggt actcccctgt 1380
getcatgect accaeagecg etgegtggae ecetggetca etcagaeceg gaagacetge 1440
cccatttgca agcagcctgt tcatcggggt cctggggacg aagaccaaga ggaagaaact 1500
caagggcaag aggagggtga tgaaggggag ccaagggacc accetgcete agaaaggacc 1560
ccacttttgg gttctagecc cactettecc acctetttg gttccttage cccagetecc 1620
cttgtttttc ctgggccttc aacagatece ccactgtece ctccctcttc ccctgttatc 1680
ctggtctaat aacccccac acatacacct ctggtgacct atttgcacag accqtcgtct 1740
tecetecagt ettetgaggg ataggggaca ttecatecea agettetece ttacceacae 1800
ctatectitt gaggggettt ggggtgggge tggggeaage agagggaetg ggtetteast 1860
tottgggcta ataaaattgt ttotttgtgg actaaaaaaa aaaa
                                                                   1904
<210> 86
<211> 1249
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incvte ID No.: 1432557CB1
cgtttatect gttgggggg gaagtgagag agecettatt cgtattgget tagatttgca 60
ageggeagtt gtctatcaaa tctatcaagt egecettage geteaggaag taegacaceg 120
```

PCT/US00/02237

```
gaaggggtgg getttgegaa gatggeggeg etgggggtge tggagteega eetgeeaagt 180
gccgtgacac ttctgaaaaa tctccaggag caagtgatgg ctgtaactgc acaagtgaaa 240
tcactgacac aaaaagttca agctggtgcc tatcctacag aaaagggtct cagcttcttg 300
gaagtgaaag accagetget geteatgtae ettatggatt tgacccacet cattetggac 360
aaageeteag gaggatetet teagggacat gatgeagttt tgagactggt agagattega 420
acggttttgg aaaagetteg teettggae caaaagetga agtateaaat tgacaagetg 480
atcaagactg cagtgacagg cagccttagt gagaatgacc cacttegttt taagcctcat 540
cccagcaata tgatgagcaa gttgagctct gaggatgagg aggaagatga agcagaagat 600
gaccagtetg aggetteagg gaagaaatet gtgaagggag tgtetaagaa atatgtteet 660
ccacgettgg ttccagtaca ttatgatgaa acagaagetg agegggagaa gaagegteta 720
gaacgagcca agagacggc attgagcagc tctgtcattc gtgaacttaa ggagcagtac 780
tcagatgetc cagaggaaat ccgtgatget cggcatcccc atgttacccg ccagagtcag 840
gaggaccaac acaggattaa ctatgaggag agcatgatgg tgcgtttgag cgtcagtaag 900
cgagagaaag gacggcgaaa acgagcaaat gtcatgagct cacaacttca ttcccttaca 960
cactteagtg acateagtgc tttgacaggg ggaactgttc atcttgatga ggatcagaat 1020
cctattaaga agcggaagaa gatacctcag aaaggtcgga agaaaaaagg ttttcggagg 1080
eggeggtgat tatgggtgta catatttgta tattttttgt catectgaga tacttctaat 1140
tteattgtat ataggtggtt tteeetggaa tteattaatt gtttgetttg gacatgtgga 1200
aagagootta otaataaaat tgattttact tatgaaaaaa aaaaaaaaa
<210> 87
<211> 1064
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incvte ID No.: 1441770CB1
<400> 87
cggctctcca gagcgtctgt aaacaccag agactgtcat ggagggggag gaggaggcgg 60
eggeggegaa gggaggegtt tggggeegee teeagggtee getetgeeat teetgaactg 120
gteectegte eeegtgaete tggeateagg gaagegaact gttaggegag aggaggagge 180
agccagaacc atateccett ettecteggg gegggggeeg ggccaggeeg getgageegg 240
gggagggete egggagggag tgeetggeea ggceggeetg tetgeegega tggatgaeag 300
taaggtggtt ggaggcaaag taaagaagcc cggtaaacgt ggtcggaagc cagccaaaat 360
tgacttgaaa gcaaaacttg agaggagccg gcagagtgca agagaatgcc gagcccgaaa 420
aaagctgaga tatcagtatt tggaagagtt ggtatccagt cgagaaagag ctatatgtgc 480
cctcagagag gaactggaaa tgtacaagca gtggtgcatg gcaatggacc aaggaaaaat 540
cccttctgaa ataaaggccc tactcactgg agaagagcag aacaaatctc agcagaactc 600
aagcaggcat accaaggctg ggaagacaga tgctaatagc aattcctggt gaagattata 660
taaagatgag tcagtgattg aagccaatat tctgattccc atggaagatg gatgggcaag 720
agtgtacttc ttggctccat ttactaccta ctgctcagta gtcatctctg taaatctgca 780
atttctacca aaatgtgtga tcgtagatct caaaggatct tgctttaact ttcaacactt 840
agaaaatcta caaacattca gacctgtctg ggttggtatt gccacccatg acatttaaca 900
tgttgtgatg cttgaaaaca caggagtaga gaaaatcgat gaagattgta tttttgcacc 960
ttaactccac attgctttat tggttaattt atattctttc catgtaattc atgtaattgt 1020
atgtetgtgt gtgttttatg tgtcaccacc tttcatgttt ttga
                                                                    1064
<210> 88
<211> 1398
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 1456684CB1
<400> 88
```

77/91

agacccaaaa gcattaagga gaaaaagaaa actacatcac ataccagggg agaaataccg 60

PCT/US00/02237

```
gaggagtcaa actatgttgc tgatcctgga ggatcactga gcaaaaccac aaatattgct 120
gaagaacca gcaaaattga aacctacatt gcaaaacctg ctctgccggg aacctccaca 180
aatagtaatg ttgcacccct ttgccaaata acagtgaaaa ttggaaacga agccattgtg 240
aaaaggcaca ttctaggatc taaattgttt tataaaagag ggagaagacc caagtatcag 300
atgraggagg agrettinge araggggaat garcragaar cragtggaga ragreracte 360
gggctttgcc aatccgagtq catggagatg agtgaagtgt tcgatgacgc aagtgaccag 420
gattecactg acaaaccgtg gegeeettac tacaactaca aacccaaaaa gaaatecaga 480
cagttgaaaa aaatgaggaa agtcaactgg aggaaggagc acggaaacag gagcccgagc 540
cataaatgta aatacccagc agaactggat tgcgccgtgg ggaaggctcc tcaggataaa 600
ccctttgagg aagaagaac taaagagatg cccaagctgc agtgtgaact ctgtgatgga 660
gacaaagcag tgggggctgg aaaccaagga aggccccacc gacatettac tteteggcca 720
tatgeetgeg agetetgege caageagtte cagageeett ccacacteaa aatgeacatg 780
agatgtcaca ccggggagaa gccataccag tgcaagacct gcggacggtg cttttcggtg 840 caaggaaact tacagaaaca tgaacgcatc cacctgggct tgaaggagtt cgtctgtcag 900
tattgcaaca aggcattcac cttgaatgag accetcaaaa tccatgaaag aatccatact 960
ggagaaaagc gttaccactg tcagttctgc tttcagagat ttttgtatct ctccaccaaa 1020
aggaatcacg agcagaggca tattcgggag cataatggga agggctatgc ctgcttccag 1080
tgccccaaaa tttgcaaaac agctgctgcc cttggaatgc accaaaagaa acacttattc 1140
aaaaqcccaa qtcaqcaqqa qaaaataqqt qacqtgtgcc acgaaaactc aaatcccttg 1200
gagaatcaac atttcattgg ttcagaagac aatgaccaaa aggataacat acaaaccggt 1260
gtggaaaatg ttgtcctttg agtggcaaga attagaaaaa tcttcaaaaa tatagttggt 1320
ggtttttttta gttatgattt aagtttagtt tcattttgtc catgtgacag tcatgaagga 1380
gtgaaaaaa aaaaaaaa
<210> 89
<211> 746
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 1602916CB1
<400> 89
eggetegagg egtetttatg ggeeceettt aaggeeggeg gaggeatete gggeegggeg 60
eggegeteeg teegteggee gtagegaetg aactgegege ggateeetee geggggetee 120
tegteccegt caegetgact tteegtgeag tgeegtggtg egaaaatgee tegeeggtge 180
gcaccggaga cagccgattt ttacgaccca gcaagaggcc gagctggtac aatatcctga 240
ctgtaaatcg tccagtggta atattggcga ggacccagac cacttaaatc agagctcgtc 300
tccttctcaa atgtttccgt ggatgagacc acaagcagct cctggtagac gaagaggaag 360
acaaacetac agtcqcttcc aaactctaga gttggaaaag gaatttettt ttaacccta 420
tetgaceagg aaaagaagaa tegaggttte eeacgeecta geecteaceg agagacaggt 480
aaaaatctgg ttccagaaca ggagaatgaa atggaaaaaag gaaaacaaca aggacaaatt 540
tecegtttee eggeaggagg tgaaggaegg ggaaacgaaa aaggaageee aagagetgga 600
ggaagacaga gccgaacgct tgacaaatta acttctacct ttaaaattta ccacagacta 660
ttaaaactaa taatcaccat atgctgtgga caccacctat tttctttgtt ggaaaagacc 720
ttactgtgtt tcaagctacc ttcatg
                                                                    746
<210> 90
<211> 1270
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 1692816CB1
gttggtcacg tggttgttcg gagcggcga gcggagttag cagggcttta ctgcagagcg 60
cgccgggcac tccagcgacc gtggggatca gcgtaggtga gctgtggcct tttgcgaggt 120
```

78/91

. Sa il Assiliante di Salatego di Liggi della segli della Migra di Maria.

PCT/US00/02237

```
getgeageea tagetaegtg egttegetae gaggattgag egteteeace eagtaagtgg 180
geaagaggcg geaggaagtg qqtacqeaqq qqcgcaaggc gcacagcete tagacgacte 240
gettteecte eggecaacet etgaageege gteetaettt gacagetgea gggeegegge 300
ctggtcttct gtgcttcacc atctacataa tgaatcccag tatgaagcag aaacaagaag 360
aaatcaaaga gaatataaag aatagttotg toccaagaag aactotgaag atgattoago 420
cttetgeate tggatetett gttggaagag aaaatgaget gteegeagge ttgteeaaaa 480
ggaaacatcg gaatgaccac ttaacatcta caacttccag ccctggggtt attgtcccag 540
aatctagtga aaataaaaat cttgqaqqag tcacccaqqa qtcatttgat cttatgatta 600
aagaaaatcc atcctctcaq tattqqaaqq aaqtqqcaga aaaacqqaqa aaqqcqctgt 660
atgaagcact taaggaaaat qagaaacttc ataaagaaat tgaacaaaag gacaatgaaa 720
ttgcccgcct gaaaaaggag aataaagaac tggcagaagt agcagaacat gtacagtata 780
tggcagaget aatagagaga etgaatggtg aacetetgga taattttgaa teactggata 840
atcaggaatt tgattetgaa gaagaaactg ttgaggatte tetagtggaa gacteagaaa 900
ttggcacgtg tgctgaagga actgtatctt cctctacgga tgcaaagcca tgtatatgaa 960
atgcattaat atttgactgt tgagaatttt actgccgaag tttacctcca ctagttcttt 1020
gtagcagagt acataactac ataatgccaa ctctggaatc aaatttcctt gtttgaatcc 1080
tgggacccta ttgcattaaa qtacaaatac tatgtatttt taatctatqa tqqtttatgt 1140
gaataggatt ttctcaqttq tcaqccatqa cttatqttta ttactaaata aacttcaaac 1200
teetgttgaa cattgtgtat aacttagaat aatgaaatat aaggagtatg tgtagaaaaa 1260
<210> 91
<211> 943
<212> DNA
<213> Homo sapiens
<221> misc-feature
<223> Incvte ID No.: 1968191CB1
<400> 91
gteccattte cagaaatcac aaggatgtta ggcaatgaat ggagtaaact gcetcetgag 60
gaaaaacagc getacettga tgaagcagac agagataagg agcgttacat gaaggaactg 120
gaacaqtatc aqaaaacaqa qqcctacaaq qtcttcaqta qqaaaaccca qqaccqtcaq 180
aaaggcaaat ctcataggca agatgcagcc cggcaggcca ctcatgatca tgagaaagaa 240
acagaggtaa aggaacggte tgtttttgac atccctatat ttacagagga attcttgaac 300
catagcaaag ctcgggaagc agagctccgc cagcttcgca aatccaacat ggagtttgag 360
gagaggaatg cagccctgca aaagcacgtg gagagcatgc gcacagcagt ggagaagctg 420 gaggtggatg tgatccagga gcggagccgc aacacagtct tacagcagca cctggagacc 480
ctgcggcagg tgctgaccag cagetttgcc agcatgccct tgcctggaag tggagagaca 540
cctacagtgg acaccattga ctcatatatg aacagactgc acagtattat tttagctaat 600
ccccaagaca atgaaaactt catagctaca gttcgagaag ttgtgaacag actcgatcgt 660
tagggaatgg tgagtgctca ctgataaata tttatatgcc agcacatcat caaaaataag 720
atgtcatcag actitatcaa tactactaaa accetgggat tacattggat gaacaagttg 780
gagacttggt taagatteet gttgcatggt tgttaaatgt agtaaataat attagaaaag 840
agaatcactg tagtcccagc tacttgggag gctgaggtgg gaggattget tgagcccagc 900
                                                                    943
agttcaagtc cagagagatc ctgtctctaa aaataaaaga aaa
<210> 92
<211> 1997
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2052061CB1
<400> 92
egaccaegeg teegetegeg tggggegega ggtacceegt ggggeageta ggtgeeteca 60
agaaaccccg ccccagagga cccgcacgag ttgttgccat ttttcgctga agcccctgcc 120
```

79/91

The or I have been not been some in the wall a property of a some of

PCT/US00/02237

```
ctgggggtgg tgttcttctc tatgattcta ccaatcgtcg gaatttttt cctcccttct 180
ttetttttag gaaggegetg gggtgtgegg eggaaaegge ggeggtgaag ggateetett 240
gtggtatctc ctccccggag aaatagggga gtgggggcc aagaacgaga agacgagaac 300
gegtegeeet gegetatgte agaatgggge gggtgtgagg ggaacagete tettgegate 360
agatcaggag tatgagcete eeggaggaeg geatgagtte tggacaette aggagteete 420
agctagtgac atggtcgata tggataaact cataaacaac ttggaggtcc aacttaattc 480
agaaggtggc tcaatgcagg tattcaagca ggtcactgct tctgttcgga acagagatcc 540
ccctgagata gaatacacaa qtaatatgac ttctccaaca ctcctggatg ccaaccccat 600
ggagaaccca gcactgttta atgacatcaa gattgagccc ccagaagaac ttttggctag 660
tgatttcage etgececaag tggaaccagt tgacetetee tttcacaage ccaaggetee 720
tetecageet getageatge tacaagetee aatacgteee eccaageeae agtettetee 780
ccagaccett gtggtgtcca cgtcaacatc tgacatgage acttcageaa acattcctac 840
tgttctgacc ccaggctctg tcctgacctc ctctcagagc actggtagcc agcagatctt 900
acatgtcatt cacactatcc cctcagtcag tctgccaaat aagatgggtg ggctgaagac 960
cateceagtg gtagtgeagt etetgeeeat ggtgtatact actttgeetg cagatggggg 1020
gcctgcagcc attacagtcc cactcattgg gggagatggt aaaaatgetg gatcagtgaa 1080
agttgaccc acctccatgt ctccactgga aattccaagt gacagtgagg agagtacaat 1140
tgagagtgga tcctcagcct tgcagagtct gcagggacta cagcaagaac cagcagcaat 1200
ggcccaaatg cagggagaag agtcgcttga cttgaagaga agacggattc accaatgtga 1260
ctttgcagga tgcagcaaag tgtacaccaa aagctctcac ctgaaagctc accgcagaat 1320
ccatacagga gagaagcctt ataaatgcac ctgggatggc tgctcctgga aatttgctcg 1380
ctcagatgag ctcactcgcc atttccgcaa gcacacaggc atcaagcctt ttcggtgcac 1440
agactgcaac cgcagctttt ctcgttctga ccacctgtcc ctgcatcgcc gtcgccatga 1500
caccatgtga geogracagg teacactaga gaagetgege tggtatettt cetggtegtg 1560
tgctgaggtt gggacaattt tttcctcttt gacttcagct tgcatatggg gttgaagcag 1620
cccactgage caagttgagg agactggagg aaaagagage tggteteeeg tggggetett 1680
catattetac etecaettet ceaetgteca gaceegtttt ttteaaeeee caeatgggtt 1740
gacttccage gtggcaccca tgggtgcctt cccatccccc cctgttctga aatagggaat 1800
ttttccccca gcaacaccac aggaatcaaa ctcaaggctg gcaaccacat ccgctgtttc 1860
ttcctcccac ttccctcttg tctctagaac tcttatccaa tgtcttaaca tcctaccaaa 1920
agggetegat gegtteeaca gattttteac ttettattge agggatacat tegtggtege 1980
cacataaggg ttgcttc
<210> 93
<211> 4334
<212> DNA
<213> Homo sapiens
<2205
<221> misc-feature
<223> Incyte ID No.: 2056207CB1
cggtggtggt ggcgggccgg ggcatgagca ggaggaggat taccgctacg aggtgctcac 60
ggccgagcag attctacaac acatggtgga atgtatccgg gaggtcaacg aggtcatcca 120
gaatccagca actatcacaa gaatactcct tagccacttc aattgggata aagagaagct 180
aatggaaagg tactttgatg gaaacctgga gaagctcttt gctgagtgtc atgtaattaa 240
tccaagtaaa aagtetegaa caegecagat gaatacaagg tcatcagcac aggatatgcc 300
ttgtcagatc tgctacttga actaccctaa ctcgtatttc actggccttg aatgtggaca 360
taagttttgt atgcagtgct ggagtgaata tttaactacc aaaataatgg aagaaggcat 420
gggtcagact attroqtqtc ctqctcatqq ttqtqatatc ttagtqqatq acaacacagt 480
tatgcgcctg atcacagatt caaaagttaa attaaagtat cagcatttaa taacaaatag 540
ctttgtagag tgcaatcgac tgttaaagtg gtgtcctgcc ccagattgcc accatgttgt 600 taaagtccaa tatcctgatg ctaaacctgt tcgctgcaaa tgtgggegcc aattttgctt 660
taactgtgga gaaaattggc atgatcctgt taaatgtaag tggttaaaga aatggattaa 720
aaagtgtgat gatgacagtg aaacctccaa ttggattgca gccaacacaa aggaatgtcc 780
caaatgccat gtcacaattg agaaggatgg tggttgtaat cacatggtct gtcgtaacca 840
gaattgtaaa gcagagtttt gctgggtgtg tcttggccca tgggaaccac atggatctgc 900
ctggtacaac tgtaaccgct ataatgaaga tgatgcaaag gcagcaagag atgcacagga 960
gegatetagg geagecetge agaggtacet gttctactgt aategetata tgaaceacat 1020
geagageetg egetttgage acaaactata tgeteaggtg aaacagaaaa tggaggagat 1080
```

80/91

geageageae aacatgieet ggattgaggt geagtieetg aagaaggeag tigatgieet 1140

Sinter and the say in the say of the same of the same

PCT/US00/02237

```
ctgccagtgt cgtgccacac tcatgtacac ttatgtcttc gctttctacc tcaaaaagaa 1200
taaccagtcc attatetttg agaataacca agcagateta gagaatgcca cagaggtget 1260
ctcgggctac cttgaacgag atatttccca agattctctg caggatataa agcagaaagt 1320
acaagacaag tacagatact gtgagagtcg acgaagggtt ttgttacagc atgtgcatga 1380
aggetatgaa aaagatetgt gggagtacat tgaggaetga gaatggeeet geataaaatg 1440
aggaggeact aagcetatte tgacaceact ggtetgtagt accagaattg ttttgttaat 1560
ggaaagttta agtaaattat attgtaataa aaaggtagat aaaccattgt agaacagtat 1620
tetaggeege caacaaaagt gtgacagaca cactaaaage cetecaactt taacttgtaa 1680
cgtagcttca ttctcaaagc tgactccttt tttttctttt tccttttcct gagtgtagta 1740
cagttaaaat ttcaaacagc tccttgacac tgcttttcat gttcaaacca gccattttgt 1800
tgtactttgg taaaggacct cttccccttc ctcccctaca catacagata cacccacaca 1860
cagactgact ctctttctct cataccccaa ggtcatgagt gaatgatgct tagttccttg 1920
taaagaaaat cttgggatgg ggaaaggggt aggcagcaag aggattcaac aaacgaaaaa 1980
cataaaaaact ttgtatatga cttttaaaac aagaggacaa cacagtattt ttcaaaattg 2040
tatatagege atatgeatgg acaaageaag egtggcacgt gtttgcataa tgtttaatta 2100
caaaaaaata tttattetti aaaaatette aagattatgt etatttgetg tgeattttet 2160
ttcagtttgc ttatctttcc cgggttgggg ttgggataaa ggtgtgtcgg tttagcacct 2220 ctggaagacc tatctagagc tetttcactt tcctqaggtt attttgccct ttctggtgtt 2280
ggtatgtetg ttgccgcca tgggcctcat gccttgaatt cctgctcttg atcagggaca 2340
agggaggtca agctctqact aatqccatqa cctqattaaq qqqtacaqca qqqaqttttq 2400
tigctacage teatgaatta acctgteeca acctaateec ectecatgge atcatgeete 2460
tacccaagee tttgtgtgcc catgttatgc acacagetgc aggcattett aagteceetg 2520
togcatocag togaagcatt ttaaaattte ttttactttt togtttteee ttaattoctg 2580
cttttcagat tttagttatg gctcgtctgc tcaccccttc tctacattag ggtgtcaaag 2640
agaatgtttt getttagata taaatageea tteatttagt eteagattgt gaatttaaaa 2700
tggtggatac cgaaattgct tgtgtgtgtt gctgtgggtt tggtttgaag gcaaacaccc 2760
ctagaacatg atatteccat ctagtgcatt taaatagaaa tcactgagtt tgctgctttt 2820
ttattgtcag cagataggag aattaataat gcattttagc tgtgatgtcc atttttatga 2880
aatteetaet aagagetatg ttaaaagtaa aggatggtgg tggttgtatt aactatatae 2940
ctgtttaggc cattetggct gtggtatttt tcaataggtc agcatetgta aatetgtcag 3000
ttttatacag gagtgcagag tgaactaggc aactagatta agaggtctaa atatgaaata 3060
ccagttgagg ctgaggacct cttcgtcttc ctttagatgt cttttgccta gggagtgttt 3120
accatttgtg aggcagettt gtetgetett acactgtaca tectattact ccattgggaa 3180
gtaggttcac tttcctctqq ccttttqcct aaqttaqqct ttqctqaatc aaccctactt 3240
ttccttttag aaaaggttgt tacaggagat ttactggcaa ctgttctttt cccatcaaaa 3300
atcagtgaat gittgctgag tataaatgct gcitccttaa accactigte gcittaggat 3360
caactttacc tgtacctttt ctcctttcct cccttgccac ctcaggtgca aatctgaact 3420
cagtgtotgc ticttocatt ttotogtoto totoccotot toccccatta tocatatgac 3480
attattttac ttcaaatgac agcatcaatc ttaaaaagat atacattaaa actaaggagt 3540
ttttttaaag aaagootgaa taagttoott toootggtaa otttgaaaag cagtcagagt 3600
tgctatatag atatatgtgg ctcctttaaa atgctttgtg tatgtgtggt gtttaaaaaa 3660
aaaaatctta ccagttaact ttgcagtgtc taggtttgag tgtcataaat ccacgtgttc 3720
ctgttgcaac aaatacccaa aaattgtgtg tgcacttcct aataccagtc ttcacccatg 3780
gaggaacagt getttttaga gatgetttet attteaatgt tggcatactg cetgagggta 3840
ttgcagttgt gggtgcattc cctaatttgt atgatcaaga tgaactggcc cttttctact 3900
tccaagcttt taacagatac caccatattt gacagaattc ccagagtgaa ttgcttgtgt 3960
tattagtaga ttcagtgccc ccagctggga taggcaagcc atgacagctt ccctgtttca 4020
cctacagaag tottatotga gggatotatt cacagtaage accaaggtot ccatgtottg 4080
aggtcagttt cattgtcttt tgaaaagtgc atgcttcatt tgaacaattc attcagcagc 4140
agatggactt tcagtgattt aaaataaaat tttgatccaa agctcaggac acaaaccaca 4200
gtggtaaaat tgagtagcat ataatatcag actaaattat ctgtaatttt ccacaaccca 4260
gattgtatgt gttttatgtg tgtttaaata aatatgttag atacacgtgt atacatacac 4320
ccatatacag caga
```

```
<210> 94
```

respondent to a state of the respondent of the state of the second of the second of the second of the second of

<211> 1706

<212> DNA <213> Homo sapiens

<220>

<221> misc-feature

```
<223> Incyte ID No.: 2101803CB1
ccgcgccccg ccgccaccat qaqqqccqaq qqcctcgqcg qcctggagcg cttctgcagc 60
cegggcaaag geeggggget gegggetetq cageeettee aggtqqqqqa ettqetgtte 120
tectgecegg cetatgecta egtgeteacg gteaacgage ggggeaacca etgegagtae 180
tgcttcacca ggaaagaagg attgtccaaa tgtggaagat gcaagcaggc attttactgc 240
aatgtggagt gtcagaaaga agattggccc atgcacaagc tggaatgttc tcccatggtt 300
gtttttgggg aaaactggaa teeteggag actgtaagac taacagcaag gattctggec 360
aaacagaaaa tccacccaga gagaacacct tcggaaaaat tgttagctgt gaaggagttt 420
gaatcacatc tggataagtt agacaatgag aagaaggatt tgattcagag tgacatagct 480
getetecate actittacte caageatete quattecetq acaatquiaq ceteqtagta 540
ctetttgcac aggttaactq taatqqcttc acaattqaaq atqaaqaact ttetcatttq 600
ggatcagega tattteetga tgttgeattg atgaatcata getgttgeee caatgteatt 660
gtgacctaca aagggaccct ggcagaagtc agagctgtac aggaaatcaa gccgggagag 720
gaggttttta ccagctatat tgatctcctg tacccaacgg aagatagaaa tgaccggtta 780
agagattett atttetttae etgtgagtge eaggagtgta ceaccaagga caaggataag 840
gccaaggtgg aaatccggaa gctcagcgat cccccaaagg cagaagccat ccgagacatg 900
gtcagatatg cacgcaacgt cattgaagag ttccggaggg ccaagcacta taaatcccct 960
agtgagetge tggagatetg cgageteage caggagaaga tgagetetgt gtttgaggae 1020
agtaacgtgt acatgttgca catgatgtac caggccatgg gtgtctgctt gtacatgcag 1080
gactgggaag gagccctqca atatggacag aaaatcatta agccctacag taaqcactat 1140
cettegtact ceetcaacgt ggeetecatg tggttgaage tagggagact ctacatggge 1200
ctggaacaca aagccgcagg ggagaaagcc ctgaagaagg ccattgcaat catggaagta 1260
getcaeggca aagateatee atatatttet qaqateaaac aqqaaattqa aagecaetga 1320
aactatgcag catttcagtt ttcatttaaa cacttagttc agaaacctta aaggatttga 1380
atatttcaaa ttgcacacgt cactccagca tctctgtaaa ataattggaa tgaaaatact 1440
tettgeactt aaacactgea catgeegtae tttgaggtta gtetgaatet tgaactttaa 1500
taccaaatta attttgaatg cttttgtttc ctaagagata atggcatggt ttcatatgtt 1560
atactttgga cagacagagt tttaaaaatg gaattatttt ttctttcatg cctcttgtaa 1620
tgttctgaac aaacttgaat gatgaaagta ttaaagagat atcagtaaaa agaacaaaaa 1680
ataaagatcc agaaagaaaa aaaggg
                                                                  1706
<210> 95
<211> 1602
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2112362CB1
<400 . 95
ccgtggggca gtcgaggatg tcggtgaatt acgcggcggg gctgtcgccg tacgcggaca 60
agggcaagtg cggcctcccg gagatetteg acccccgga ggagctggag cggaaggtgt 120
gggaactggc gaggctggtc tggcagtctt ccaatgtggt gttccacacg ggtgccggca 180
tcagcactgc ctctggcatc cccgacttca ggggtcccca cggagtctgg accatggagg 240
agegaggtet ggececaag ttegacacca cetttgagag egegeggec acgeagacc 300
acatggcgct ggtgcagctg gagcgcgtgg gcctcctccg cttcctggtc agccagaacg 360
tggacgggct ccatgtgcgc tcaggcttcc ccagggacaa actggcagag ctccacggga 420
acatgtttgt ggaagaatgt gccaagtgta agacgcagta cgtccgagac acagtcgtgg 480
geaccatggg cetgaaggee aegggeegge tetgeacegt ggetaaggea agggggetge 540
gagectgeag gggagagetg agggaeacea tectagaetg ggaggaetee etgeeegaee 600
gggacctggc actogccqat qaggccaqca qqaacqccqa cctqtccatc acqctqqgta 660
categorigea garceggee agegggaace tgeegergge taccaagege eggggaggee 720
geetggteat cgtcaacctg cageccacca ageacgaeeg ccatgetgae etcegeatec 780
atggctacgt tgacgaggtc atgacccgc tcatgaagca cctggggctg gagatccccg 840
cetgggacgg cccccgtgtg etggagaggg cgctgccacc cctgccccgc ccgcccaccc 900
ccaagetgga geccaaggag gaateteeca eeeggateaa eggetetate eeegeeggee 960
ccaagcagga geeetgegee cagcacaacg getcagagee egecageeee aaacgggage 1020
ggcccaccag ccctqcccc cacaqaccc ccaaaaqqqt gaaqqccaaq qcqqtccca 1080
gctgaccagg gtgcttgggg agggtggggc tttttgtaga aactgtggat tcttttctc 1140
```

PCT/US00/02237

```
tegtggtete actitgtiae tigtitetgt eeeegggage eteagggete igagagetgt 1200
getceaggee aggggttaca cetgeeetee gtggteeete cetgggetee aggggeetet 1260
ggtgcggttc cgggaagaag ccacaccca gaggtgacag ctgagccct gccacaccc 1320
agectetgae ttgetgtgtt gtecagaggt gaggetggge cetecetggt etceagetta 1380 aacaggagtg aacteeetet gtececaggg cetecettet gggeeeceta cageccace 1440
taccetect ccatgggee tgeaqqaqqq qagacecace ttgaagtqqq ggateagtag 1500
aggettgeac tgeetttggg getggagga gacgtgggte caccaggett etggaaaagt 1560
cctcaatgca ataaaaacaa tttctttctt gcaaaaaaaa aa
<210> 96
<211> 1951
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2117346CB1
<400> 96
gaggegege egacegege etetttegeg eggattaggg ggteteggeg agggagteat 60
caagetttgg tgtatgtgtt ggeeggttet gaagtettga agaagetetg etgaggaaga 120
ccaaagcagc actogttgcc aattagggaa tggaccgttt gggttccttt agcaatgatc 180
cototgataa gocacettge egaggetget ectectacet catggagget tatatcaagt 240
gtgctgaatg tgggccacct cettttttcc tetgettgca gtgtttcact cgaggetttg 300
agtacaagaa acatcaaagc gatcatactt atgaaataat gacttcagat tttcctgtcc 360
ttgateccag etggaetget caagaagaaa tggeeetttt agaagetgtg atggaetgtg 420
gctttggaaa ttggcaggat gtagccaatc aaatgtgcac caagaccaag gaggagtgtg 480
agaagcacta tatqaaqcat ttcatcaata accetetgtt tgcatctace etgetgaace 540
tgaaacaagc agaggaagca aaaactgctg acacagccat tccatttcac tctacagatg 600
accetececg acetacettt gaeteettge ttteteggga catggeeggg tacatgeeag 660
ctegageaga ttteattgag gaatttgaca attatgeaga atgggacttg agagacattg 720
attituttga agatgactcg gacattttac atgctctgaa gatggctgtg gtagatatct 780
atcattccag gitaaaggag agacaaagac gaaaaaaaat tataagagac catggattaa 840
teaacettag aaagttteaa ttaatggaac ggeggtatee caaggaggte caggacetgt 900
atgaaacaat gaggcgattt gcaagaattg tggggccagt ggaacatgac aaattcattg 960
aaagccatgc attqqaattt gaactccgaa gggaaatcaa gaggctccaa gaatacagga 1020
cagcaggcat taccaatttt tgtagtgcca gaacctacga tcacctcaag aagacacggg 1080
aggaagageg cettaaaege actatgetet cagaagttet ceagtatate caggacagta 1140
gtgettgeea geagtggete eqeeqqeaaq etqacattga tteeggeetg agteetteea 1200
ttccaatggc ttcgaattca qqtaqacqqa qtqcaccacc cttgaacctc actqqcctcc 1260
ctggcacaga gaagetgaat gaaaaagaaa aggagetetg teagatggtg aggttggtee 1320
ctggagecta tttagaatac aaatetgete tattgaacga atgtaacaag caaggagget 1380
taagactggc gcaggcaaga gcactcatca agatagatgt gaacaaaacc cggaaaatct 1440
atgatttcct catcagagaa ggatacatca ctaaaggcta aggctccaag agcttgggat 1500
cagaagtcag aagtttggaa tgtggtgggt caaaggacaa tatgggtggg cattctggag 1560
agttgttttt cagctgaatt ctcatggtga aaacagggga aaggacaaag gaaaccttaa 1620
gttgtattgt ctactttett etceatectg etttaaaaca etcetgttgt tggtattatg 1680
ctgcagagtt gtgtqctaca taaqctatta ttaaatgtga gtgggcattc attcctaaca 1740
ctaaacccac agtotgtaaa toccaggagt toaagcctag aatoottaat attgtooggg 1860
gaaaggtttt tctgaataaa acacggcttt ggcctttaaa aaaactttgg ggaaacgaat 1920
tttaaaaatt aaaagggaaa agggttttt a
<210> 97
<211> 854
```

<212> DNA

<213> Homo sapiens

<220>

· 克拉克·西拉拉斯 (1)

83/91

WO 00/44900 PCT/US00/02237

```
<221> misc-feature
<223> Incyte ID No.: 2119917CB1
<400× 97
cccacgcgtc cgcggacggt gggcgcgcgc tatgacggcc agcgcacagc cgcggggcg 60
geggecagga gteggagteg gagtegtggt gaccagetge aageateege gttgegteet 120
cctggggaag aggaaagget cggttggage tggcagtttc caactecctg gaggtcatct 180
ggagtteggt gaaacetggg aagaatgtge teaaagggaa acetgggaag aageagetet 240
tcacctgaaa aatqttcact ttqcctcaqt tqtqaattct ttcattqaga aggagaatta 300
ccattatgtt actatattaa tgaaaggaga agtggatgtg actcatgatt cagaaccaaa 360
gaatgtagag cctgaaaaaa atgaaagttg ggagtgggtt ccttgggaag aactacctcc 420
cctggaccag cttttctggg gactgcgttg tttaaaagaa caaggctatg atccatttaa 480
agaagatctg aaccatctgg tgggatacaa aggaaatcat ctctaggtgg ccgagaagat 540
ttgattttct ttaaaaagac aagaataagg tctggttagg gaatgaaaaa tgtatacatt 600 tcggaacaac tccattttat ctaaaaaagt tcttgtgatt gccagtttat ttgcagtctc 660
ttaatgtate ecceatett teagecagta ettgagaaaa tttttetgaa atatgteatt 720
gaattgtatt ccaqacacaq aatacatgat aaatactgat attatgggta atctgctttc 780
catatttacc tatggatatg tacgtgcaat gtgccataac tagttgagag ggtgggtgag 840
gcagattttc ctrr
<210> 98
<211> 1581
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2123456CB1
teegggaaag tttetttgga ggteeggeee ggageggeea tgteecaegg eeceaageag 60
cccggcgcgg ccqccqtqcc qqcqqqcqc aaqqctccgg gccagcatgg gggcttcgtg 120
gtgaetgtca agcaagagcg eggegagggt ecaegegegg gegagaaggg gteceaegag 180
gaggageceg gttetgeage eggtgaagaa aegeggetgg eecaagggea agaageggaa 240
gaagattetg eegaatggge eeaaggeace ggteaeggge taegtgeget teetgaacga 300
geggegegag cagateegea egegeeacce ggatetgeee ttteeegaga teaccaagat 360
getgggegee gagtggagea agetgeagee aaeggaaaag eageggtace tggatgagge 420
cgagagagag aagcagcagt acatgaagga gctgcgggcg taccagcagt ctgaagccta 480
taagatgtgc acggagaaga tccaggagaa gaagatcaag aaagaagact cgagctctgg 540
getcatgaac acteteetga atggacacaa gggtggggac tgcgatgget tetecacett 600
cgatgttccc atcttcactg aagagttctt ggaccaaaac aaagcgcgtg aggcggagct 660
teggegettg eggaagatga atgtggeett egaggageag aacgeggtae tgeagaggea 720
cacgcagage atgageageg egegegageg tetggageag gagetggege tggaggageg 780
gaggacgetg gegetgeage ageageteea ggeegtgege caggegetea eegeeagett 840
cgcctcactg ccggtgccgg gcacgggcga aacgcccacg ctgggcactc tggacttcta 900
catggeeegg etteaeggag ceategageg egaceegee eageaegaga ageteategt 960
cegeateaag gaaateetgg eecaggtege cagegageae etgtgaggag tgggegggee 1020
cacgatgcag aggagaagct gtgggcgcgg ccctgccaca ccccaccccg tggacgagag 1080
getgggggte caccetttgg ggeetggtee cateetgeae ettggggget ceageecece 1140
taaaattaaa tttetgeage ateeetttag ettteaatet eeceageece etgaaceegg 1200
aaaaagcact cgctgcgcga tacacccaga agaacctcac agccgagggt gcccctcctc 1260
ggaggacage caegegetae actggetete egggecaece ceaggacaea gggcagaega 1320
aacccacccc cagcacacgg caggaccccc caaattactc actacggggg gctgtgccat 1380
aggecacaca ggaagetgee ttgtggggae ttacetgggg tgteeceege atgeetgtae 1440
cccagatggg tgggggccgg ctttgcccat cctgctctcc tccagccgag ggaccctggt 1500
gggggtggct ccttctcact gctggatccg gactttttaa ataaaaacaa gtaaaatttg 1560
tgttttaaaa aaaaaaaaa a
                                                                    1581
```

<210> 99 <211> 2150

PCT/US00/02237

WO 00/44900

```
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incvte ID No.: 2148792CB1
gtotogatot cotgacotog coatocacco gootcagoca cocgaagtgt tgggatttca 60
ggggccaaat agttgcatca catgtatcta atccgagagt ctcatgcttc tggtagctcc 120
tcagtgacca getcetgete actgecetea gaaageeeaa ceetcaggea atggeggeet 180
tgttcctgtc tgccccaccc caggccgagg tgaccttcga ggacgtggct gtgtacctct 240
cccgggagga atggggccgc ctgggccctg ctcagagggg cctctacagg gacgtgatgc 300
tggagaceta egggaaceta gteteaetgg gagtaggaee tgeaggeeec aageetggag 360
tgatetegea gttggagega ggggatgage cetgggteet ggatgtteag ggcacetetg 420
ggaaagagca cctgagagtc aacagcccag ctcttgggac cagaactgag tacaaggagt 480
tgacttcaca ggagacattt ggtgaggaag atccccaggg atctgagcca gtagaagcct 540
gtgaccacat cagtaagtca gaggggagcc tggaaaagct agtggagcag agaggcccca 600
gggcagtcac actgaccaac ggggagagca gcagggagtc tgggggaaac ctcaggttgc 660
tgtcaagacc tgttcctgat cagagacctc acaaatgtga tatatgtgag caaagttttg 720
aacagagatc atatctcaac aaccataagc gtgtacacag gtcaaaaaaa acaaatacag 780
ttcgtaactc tggggaaatc ttcagtgcaa acttagttgt taaagaagat cagaaaattc 840
ctactgggaa aaaattgcat tattgcagtt actgtgggaa aacattcagg tacagtgcca 900
accttgtcaa gcatcagcgg cttcacactg aagagaagcc ctacaaatgt gatgagtgtg 960
ggaaageett cageeagage tgegagttea teaatcaceg aaggatgeae teaggagaga 1020
ttccctaccg gtgtgacgag tgtgggaaga cattcacccg gaggcccaac ctcatgaage 1080
accagaggat tcacactggg gagaaaccct acaagtgtgg ggagtgtggg aagcacttta 1140
gegeetacte tteeetgatt tateaccaga gaatecacae eggagagaaa ecetataaat 1200
gtaatgactg cgggaaagcc ttcagtgatg gctcaatcct tatccgacat cgtcggactc 1260
acaccggaga gaagccattt gagtgcaagg aatgtggcaa aggctttaca caaagttcta 1320
accttateca acateagaga atteacactg gagagaaacc ctataaatgt aatgaatgtg 1380
agaaagettt cattcaaaaa accaaacteg tggaacatca gagaagecac actggagaga 1440
agccctatga atgcaatgac tgtggcaaag ttttcagcca aagcacacac ctcatccagc 1500
accagagaat ccacacagga gagaagccct acaagtgcag cgagtgtggg aaagccttcc 1560
acaacagtte cagacteate caccaccaga ggetgeacca eggagagaaa ecetacagat 1620
geagegattg caagaaagee tteageeaga geaegtaett gatteageae eggaggatee 1680
acaccgggga gaagccctac aagtgcagcg agtgtgggaa ggccttccgg cacagttcca 1740
acatgtgtca gcatcagcgg attcacctcc gggaggactt ctccatgtaa cagtggcgcg 1800
gtgtccgagg gcagagtcca gctgagcact tcctgcatgc gcccccggca cctgactctg 1860
ccctttatgt attatccaca cgatgttttc acagagtgaa aggacgtttc tcattaaaca 1920
aacetettt ettaaateaa aaaaaaaaa agggggggee getttagggg teecaggggt 1980
toccaaccco taatoctaaa caatgtaata gotgttocog gtgtaaaatt gttagoggco 2040
caaaattccc tggcaaatta tcgaaccggg aatccttaaa tgtttaaaac ccccggggtg 2100
                                                                    2150
geecceaaaa agggttgaet egaaaettee acattaaaat tegggggggg
<210> 100
<211> 691
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 2751943CB1
<400> 100
eggegeegga gegggegtea tggegegget cetetggttg eteeggggee tgaeeetegg
aactgcgcct cggcgggcgg tgcggggcca agcgggcggc ggcgggcccg gcaccgggcc 120
gggactgggg gaggcagggt ctcttgcaac gtgtgagctg cctcttgcca agagtgagtg 180
gcaaaagaaa ctaaccccgg agcagttcta cgtcacaaga gaaaagggaa cggaaccgcc 240
tttcagtggg atctacctga ataacaagga agcaggaatg tatcattgcg tgtgctgcga 300
cagtccacte tteagttetg agaaaaagta etgetetgge actgggtgge ettegtttte 360
cgaggeteat ggtacgtetg getetgatga aageeacaca gggateetga gaegtetgga 420
```

85/91

THE HEAT WALL TO THE THE THE

PCT/US00/02237

```
tacctcgtta qqatcaqctc qcacaqaqqt tqtctgcaag cagtgtgaaq ctcatctagg 480
tcacgtgttt cctgatggac ctgggcccaa tggtcagagg ttttgcatca acagtgtggc 540
tttgaagttc aaaccaagga aacactgacc atcttcaaga gtcccgttcc cttgccaccc 600
cttcacqtqc accctcaatt tccacaattc acttqaatqa cttgttttat ttqcaataaa 660
actgggctga atttgcaaaa aaaaaaaaaa a
<210> 101
<211> 2101
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incvte ID No.: 3128913CB1
<400> 101
gtggcttgca getcggggtg ggtggctcat tteetggeeg etcetggget tegeggaaag 60
aagagattac tcacactcet tctcaagcac agaaccagtt gtactgagct ttttgctaag 120
ctgtttcagc caagaatggc tgtggaatct ggagtgattt caaccctgat acctcaggat 180
cctccggaac aagaactaat actagtgaaa gtagaagata acttttcctg ggatgagaaa 240
tttaagcaga atgggagtac tcaatcctgc caagaattgt ttcgtcagca attcagaaaa 300
ttttgctacc aggagacacc tgggccccgg gaggctctga gccgactcca ggaactttgc 360
tatcagtggc taatgccaga gttgcacaca aaggagcaga tettagaact getggtactg 420
gagcagtice tgagcattet geetgaggag etgeagatet gggtteagea acataateca 480
gaaagcggcg aggaagctgt gaccctgttg gaggatttag agagggagtt tgatgacca 540
gggcagcagg tcccagctag tccacaggga ccagcagtgc catggaagga tttaacatgt 600
ctcagagcat cccaagagte aacagacate cacetecage cettaaagac acagetgaaa 660
tcctggaaac catgcctttc ccctaaaagt gattgtgaga acagtgaaac agcaacaaaa 720
gagggcatct cagaagaaaa atcacaggga ctccctcagg aaccttcatt tcgaggaatt 780
agtgagcatg aaagcaattt agtgtggaag caaggaagtg ctacagggga gaaactaaga 840
totoottooc aagggggeag ttttagteaa gtgatottoa caaacaaato totaggaaag 900
agagacettt atgatgagge tgaaagatge ttgattetaa etacagaete tataatgtgt 960
cagaaagtto otocagaaga gagacottat agatgtgatg tatgtgggca cagottcaag 1020
cagcatteet etetaacaca acateagaga atecatactg gagaaaagee etataaatgt 1080
aaccagtgtg ggaaggcett tagtttgagg teetatetta ttatteatea gagaatteat 1140
agtggtgaga aagcatatga atgtagtgaa tgtggggaaag ctttcaatca gagctcagcc 1200
ctcattagac atcggaaaat ccatactggt gagaaagctt gtaaatgtaa tgagtgtggc 1260
aaagcattca gtcaaagttc atatctcatt atacatcaaa gaattcacac tggtgagaaa 1320
ccttatgagt gtaatgaatg tgggaaaacc tttagccaga gctcaaaact cattagacat 1380
cagogaatte acacaggaga gagaccetat gaatgtaatg aatgtggaaa agetttcagg 1440
cagageteag agetgattae teateagaga atacatagtg gagagaaace etatgaatgt 1500
agtgaatgtg gaaaagettt cagtttgage teaaacetta teagacatea gagaatteat 1560
agtggggagg aaccttatca gtgtaatgaa tgtggcaaaa ctttcaaaag gagctcagcc 1620
cttgttcagc atcagagaat tcattctggg gatgaagctt atatatgtaa tgaatgtggg 1680
aaggetttea ggeacagate ggteettatg egecateaaa gagteeacae tataaagtaa 1740
tttgtgaata ctgtgaatag tgtaaatact tcagtcagat ttttaagttt gttagtcaaa 1800
agagtttact ttggagcaaa actccataaa ggttataaaa tactaggtct tgagtctagc 1860
ttgctttgtg cagcatttcc cagtgctaat gtaaagtgtc ccttgaaagc ttttcctgtg 1920
actaatcaga acagaataca gaagaatcat tacttccagc tcttctctta ttaggaatac 1980
tcaggaaata cgaaaagtgg gaatgtaaca ttgaaacctc attttgtatg aaagtgtcat 2040
gaatatagca acccaggete tgtcactgca tetgattgtt agtgtgccag gttatggggg 2100
                                                                    2101
<210> 102
<211> 2196
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 3282941CB1
```

And the state of the state of the state of the state of the state of

PCT/US00/02237

```
<400> 102
cctactttct ctagaccaga aaattcccaa aattgtgcac attagaatca cgtgggtagc 60
ccacaaaatt cccagtgcca aattataacc tgcctttcct gtggcccagt ggtttccaac 120
gacattgggc cctcggtctg agaagtqcct ataatactgg cttacatggc cctcaggctg 180
aaccttggac gtggcaggta ttgcaaatat tttccqcqqt gcqcgcqqat tctqqacttq 240
ggcgccaact cgtagtccac gctccccggg gtcagcagag gggcgtcacg ctctcgccac 300
ccacctoget tteteaccc gegetteceg geetgggttt ttagtettee ttggageget 360
ctctggcctc cgcctccgcc agggagegga aggcggagac agcgagactg gccagggggg 420
aggaaagagg acgcgtgtgg gcaaggggga caacgggatg tccacgggct cggtgagtga 480
teeggaggag atggagette gggggetgea gegggagtae eeggteeeeg eetecaagag 540
geogecete egeggegtag agegeageta egeetegeee agtgacaact egteggeaga 600
ggaggaggac cccgacggcg aggaggagcg ctgcgctctg ggcacagccg gcagcgcgga 660
aggetgeaag aggaagegge eeegtgtgge tgggggegge ggegeaggtg gtagegegg 720 eggtggtgge aagaageee teeeggeaa gggeteagee geaqagtge aagaageee
geggaacgeg gecaacgeec qtqaqeqtqc eegqatqeqc qtqetqaqea aageettete 840
caggeteaag accageetge cetgggtgee eccegaeaet aageteteea agetggaeae 900
geteeggetg getteeagtt acategetea cetgeggeag etgttgeagg aggacegeta 960
tgagaacggc tacgtgcacc cagtgaacct gacatggcca ttcgtggtct cgggaagacc 1020
ggactotgac accaaagaag tttoogcago caacagacta tgtggaacca cogottaaat 1080
eggactggaa etcaettgat gggattatte gttaaatgeg agtgtttggg ggecaeggag 1140
agaagggaga getegtgaga toggaagaag ttteegetgg atteteettg accetteece 1200
tttccctgga actgtgatcg tgacaggtgg cgggtgtggc tgtcactgca cagcgcccac 1260
ggctacaget gegeeggate tgggegacea eqtititgeet etecaaaaag agetteetit 1320
cgtgacgaga cgcggacgca ggtccaccct cgggccctag ctctgtagac taactctcgg 1380
ctgctgcccc agcccgcgcc agacagccca cqqatccqtt ctcaqcqqac qcaqattcat 1440
egeacacgtg egggaeggtt ecacacagee ecqqeettte geggtgacae aatqqttagq 1500
gaacggttag aacgcgctct acatccgctt taaaqacaqa ggtctaqacg tgagatccgc 1560
gtcgggacag ggttttaagt gacaaagaag ggcgagtggc ttctctgggc cgggttcgta 1620
ctccagcaca gegecettet aacgggeggg aggaaggeeg etgetegeag ggetaggtgg 1680
agacacactt cocagatoac cgcaggcggg ttttacccgg agagetetgg geegttegge 1740
ctccctgccg ggtggcttct tcaatcccgt ctccttccca agctcccggc tttttctaat 1800
caggeaggeg tetgteaace etetecaett etgggetgaa geeteecaa geeeggtge 1860
gecaacetgt gtggggtett ettegggeet eesteteege eeegeteetg etectaeetg 1920
cagcaccccc agctccgact ccagactctc tgcatcaggt ctccccactc cacgctccgg 1980
gegeeceaac tecaacacca egteetgeeg egeaggttet teecegegeg gaggagegeg 2040
cagggtgggc ggcttaccat agcaagtgat cctgcgatag ggaacgcgcc cttgccccga 2100
ggctgcacta ccacaggaaa taacatatgt aaataaattt atttttttat gaataataaa 2160
acgcgctgta aaaaccgtga aaaaaaaaa aaaagg
                                                                   2196
<210> 103
<211> 749
<212> DNA
<213> Homo sapiens
<220>
<221> unsure
<222> 735, 736, 741
<223> a or g or c or t, unknown, or other
<223> Incyte ID No.: 3286656CB1
<400> 103
getgagetgt getgegegge geggegegge geggtgegge aeggeaeggt gggagtgtet 60
coggetgget tgcagggaga acaccgactg agacctcaaa ccctggetcc agtqtcatgg 120
aatccgtcac ctttgaggat gtggccgtqq aqttcatcca qqaqtqqqca ttqctqqaca 180
gegeaeggag gageetgtge aaatacagga tgettgaeca gtgeaggaee etggeeteea 240
ggggaactec accatgcaaa cocagttgtg teteccaget ggggcaaaga gcagagccaa 300
aggcaacaga acgagggatt ctccgtgcca caggtgttgc ctgggaatct caacttaaac 360
cogaagagtt goottotatg caggatottt tggaagaagc atootocagg gacatgcaaa 420
tggggccggg gctgttcctg aggatgcagc tggtgccctc catagaagag agggagacac 480
cattgacteg agaggacegg ccaqetetee aggageegee ttggtetetg ggatgeaegg 540
gactgaagge egetatgeag atteagaggg tggtgatace agtgeetact etgggeeace 600
gcaacccatg ggtggccagg gattetgcca tgtagcacgt gcctgcttcc cctttgcctt 660
```

87/91

PCT/US00/02237

```
cogcoatgat totaagttte togagteete cocagocatg cgtcctgtac aacctgtgga 720
accaggcage caaanneagt natagttgt
                                                                     749
<210> 104
<211> 1311
<212> DNA
<213> Homo saniens
<221> unsure
<222> 1294
<223> a or g or c or t, unknown, or other
<223> Incvte ID No.: 3490802CB1
<400> 104
qqqcctttqt ttctcqctqc aqcqqqaqct ccaqqtttat cctctqtqtt ctgtqtcctg 60
tgcttataga ggcccgtcct ctgtggccgt gtgacctgca agtattggga gagccacagc 120
taaaccccgg gacccctgga agcctagaaa tgggaccatt gcaatttaga gatgtggcca 180
tagaattete tetggaggag tggcattgce tggacactgc acageggaat ttatataggg 240
atgtgatgtt agagaactac agaaacttgg tetteettgg tattgttgte tetaagecag 300
acctggttac ctgtctggag caaggaaaaa aacctttaac tatggaaaga catgagatga 360
ttgccaaacc cccagttatg agttctcatt ttgcccaaga cctttggcca gagaacatac 420
aaaattettt eeaaataggg atgetgagaa gatatgaaga atgeagaeat gacaatttae 480
agttaaaaaa aggctgtaaa agcgtgggtg agcataaggt gcacaaagga ggttataatg 540 gacttaacca atgtttgaca actacccaga aagaaatatt tcaatgtgat aaatatggaa 600
aagtetttea taagttttea aatteaaaca catataagae aagacataet qqaataaate 660
ttttcaaatg tataatatgt ggcaaagctt ttaaacggtc ctcaaccctt actacacata 720
agaaaattca tactggagag aaaccttaca aatgtgaaga atgtggcaaa gcttttaacc 780
aatcctcaaa ccttactaca cataaqaqaa ttcatactqq agagaaacct tacaaatgtg 840
aagaatgtgg caaagctttt aactggtcct cagaccttaa taaacataag aaaattcata 900
ttgaacgaaa accetacata gtgaagaatg tgacagatet tttaaatgtt cetecaettt 960
taattagcat aagataattc atactggaga gaaaccctat gaatgtgatg aatgtgggaa 1020
agcetttaac cagecetega etettagtaa atttgagagt ttatatggaa cacaaaccet 1080
acaaatataa agaatgtgac aaagcttttt aaggaagttc tcaaccctta ttacacataa 1140
ttcataccaa acagaagccc tacaagtgtg aagaatgtgg caaaacctat aaacctataa 1200
caagttetea atteetttt tittgagatg gagttteact ettgteaceg aggeagaggt 1260
tgcagtgage actccagtct aggcgacaga gtangecttg tcgcgatccc a
<210> 105
<211> 990
<212> DNA
<213> Homo sapiens
<220>
<221>
<223> Incyte ID No.: 3507366CB1
<400> 105
caaaaaggaa agaataaggc aaattottag aatgtatgca caacttaata toottgctcg
tttgatgtgc attgatctca tttaataggt ttttgaatct gaaaattcaa gaaggtgaag 120
ctcacaacat tttttgccct gcatatgatt gcttccaact tgtacctgtg gatatcatag 180
aaagtgtagt ttcaaaggag atggacaaac gatacctaca gtttgatatt aaggcctttg 240
ttgaaaataa teetqeeatt aaatqqtqte etaeteeagg etgtgacaga geagtaagac 300
taacgaaaca agggtcaaat acatetggat etgatacact cagetteeca ttgetgagag 360
ctcctgctgt tgattgtgga aaaggacacc tcttctgctg ggagtgcctt ggtgaagcac 420
atgageettg tgactgeeaa acatggaaga attggetgea aaaaataacc gaaatgaaac 480
cagaagaact tgtgggagtt agtgaagcct acgaggatgc cgccaattgt ctctggttat 540
taactaactc caagccttgt gccaactgta agtctccaat acagaagaat gaaggctgca 600
atcacatgea gtgtgctaag tgcaagtatg acttttgctg gatttgcctt gaagagtgga 660
aaaaacatag ttegteeact ggaggttatt acagatgtac tegetatgaa gteatteaac 720
acgtggagga gcaatccaag gaaatgactg tggaggctga gaaaaaacac aaacgatttc 780
```

88/91

But the winds in the same of the first of the same

PCT/US00/02237

```
aggaacttga cagatttatg cactattata caagatttaa aaaccatgag catagttatc 840
agctagaaca acgccttctt aaaacagcca aagaaaagat ggagcaaatg agcagagtct 900
caaagaactg aaggaggetg tecagatace acttteattg gagatgagtt catgtgetet 960
aaaaatcggc gcatctcaag tgtcttatca
<210> 106
<211> 1048
<212> DNA
<213> Homo sapiens
-2205
<221> misc-feature
<223> Incyte ID No.: 3573060CB1
<400> 106
ccgctgcagc tetecgcggg acateteacc gttctggaga cagggetege tegeteteac 60
ggettettag geeggggttg gacageegee tteeggeeag aggggatgag gttgegetge 120
gctccgggag cgccgatggc gtgactggcc ccgcgcggag cagcgacact gcccggccag 180
cccgettete tgcccggage catgaatete agtagegeca gtageaegga ggaaaaggea 240
gtgacgaceg tgctctgggg ctgcgagete agtcaggaga ggcggacttg gacettcaga 300
ccccagctgg aggggaagca gagctgcagg ctgttgcttc atacgatttg cttgggggag 360
aaagccaaag aggagatgca tegegtggag ateetgeeee cagcaaacca ggaggacaag 420
aagatgcago oggteaccat tgeotoacte caggootoag tectococat ggtetecatg 480
gtaggagtgc agetttetee eccagttact ttecagetee gggetggete aggaccegtg 540
ttcctcagtg gccaggaacg ttatgaagca tcagacctaa cctgggagga ggaggaggaa 600
gaagaagggg aggaggaagga agaggaagag gaagatgatg aggatgagga tgcagatata 660
tototggagg agcaaagcco tgtcaaacaa gtcaaaaggc tggtgcccca gaagcaggcg 720
agcgtggcta agaaaaaaa gctggaaaaa gaagaagagg aaataagagc cagcgttaga 780
gacaagagcc ctgtgaaaaa ggccaaagcc acagccagag ccaagaagcc aggattcaag 840
aaatgaggag ccacgccttg gggggcacgg tgcaaagtgg gccttccctg ggctgtgctg 900
caggcacagg gtgcccctgt ccagcccctc cacctgtgtc tgaatgcaac aggggtgttg 960
cgggggcaac atgagagccc ctcacccca actetecact ttcaggaggc ccccagtgaa 1020
gagececace tegggteaca ataagtgt
<210> 107
<211> 1349
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 3573661CB1
gagaaagcag agctatttca agagtgagcc acagaaggga atccagaggc catctaagcg
aggaagggtc tacaggcagt gagtgaaggc caggagcagg gcccaggcca ggcacgacca 120
ccgaggggat gaacttcaca gtgggtttca agccgctgct aggggatgca cacagcatgg 180
acaacctgga gaagcagctc atctgcccca tctgcctgga gatgttctcc aaaccagtgg 240
tgatectgee etgecaacae aacetgtgee gcaaatgtge caacgacgte ttecaggeet 300
cgaatcetet atggeagtee eggggeteea ceaetgtgte tteaggagge egttteeget 360
geccategtg caggeatgag gttgteetgg acagacaegg tgtetaegge etgeagegaa 420
acgtgctagt ggagaacatt atcgacattt acaagcagga gtcatccaag ccgctgcact 480
cgaaggctga gcagcacctc atgtgcgagg agcatgaaga agagaagatc aatatttact 540
geetgagetg tgaggtgeec acetgetete tetgeaaggt etteggtgee cacaaggaet 600
gtgaggtggc cccactgccc accatttaca aacgccagaa gagtgagctc agcgatggca 660
tegegatget ggtggeagge aatgacegeg tgcaageagt gateacacag atggaggagg 720
tgtgccagac tatcgaggac aatagccgga ggcagaagca gttgttaaac cagaggtttg 780
agageetgtg egeagtgetg gaggagegea agggtgaget getgeaggeg etggeeeggg 840
agcaagagga gaagetgeag egegteegeg geeteateeg teagtatgge gaccacetgg 900
aggeeteete taagetggtg gagtetgeea tecagteeat ggaagageea caaatggege 960
```

PCT/US00/02237

```
tgtatctcca gcaggccaag gagctgatca ataaggtcgg ggccatgtcg aaggtggage 1020
tggcagggcg gccggagcca ggctatgaga gcatggagca attcaccgta agggtggagc 1080
acgtggccga aatgctgcgg accatcgact tccagccagg cgcttccggg ggaggaagag 1140
gaggtggccc cagacggaag aagagggcaa cccgggggcc ggaagaaaaa acggcccgga 1200 tgggggcctta taggcctttg cgcccgaacc ccgacccct gcttcgaaaa agccccggc 1260
gettaagaat tteeggggga aggaattett geegeaaaaa aacceeggea agettttaac 1320
cccccaaaat tccggggcgc ccggggccc
<210> 108
<211> 1642
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 3633422CB1
<400> 108
ctagttctag atcgcgagcg cccgcccagg cacccaccag cggcgccgac ctcagcgcgc 60
acctatgggc tegetaccag qacatgegga qactggtgca egaceteetg eeeeeegagg 120
tetgeagtet cetgaacca geagecatet acgecaacaa egagateage etgegtgacg 180
ttgaggtcta cggctttgac tacgactaca ccctggccca gtatgcagac gcactgcacc 240
cegagatett cagtacegee egtgacatee tgategagea etacaagtae ecagaaggga 300
tteggaagta tgactacaac eccagetttg ceateegtgg cetecaetat gacatteaga 360
agageettet gatgaagatt gacgeettee actaegtgea getggggaca geetacaggg 420
geetecagee tgtgccagae gaggaggtga ttgagetgta tgggggtaee cageacatee 480
cactatacca gatgagtggc ttctatggca agggtcctc cattaagcag ttcatggaca 540
tetteteget aceggagatg getetgetgt cetgtgtggt ggactactit etgggecaca 600
gcctggagtt tgaccaagca catctctaca aggacgtgac ggacgccatc cgagacgtgc 660
atgtgaaggg ceteatgtae cagtggateg ageaggacat ggagaagtae ateetgagag 720
gggatgagac gtttgctgtc ctgagccgcc tggtggccca tgggaaacag ctgttcctca 780
tcaccaacag tcctttcagc ttcgtagaca aggggatgcg gcacatggtg ggtcccgatt 840
ggcqccaqct cttcqatqtq qtcattqtcc aggcaqacaa qcccaqcttc ttcactqacc 900
ggcgcaagcc tttcagaaaa ctcgatgaga agggctcact tcagtgggac cggatcaccc 960
gettggaaaa gggcaagate tateggeagg gaaacetgtt tgaettetta egettgaegg 1020
aatggcgtgg cccccgcgtg ctctacttcg gggaccacct ctatagtgat ctggcggatc 1080
teatgetgeg geaeggetgg egeaeaggeg ceateatece egagetggag egtgagatee 1140
gcatcatcaa cacggagcag tacatgcact cgctgacgtg gcagcaggcg ctcacggggc 1200
tgctggageg catgcagace tatcaggacg eggagtegag gcaggtgetg gctgcctgga 1260
tgaaagageg geaggagetg aggtgeatea eeaaggeeet gtteaatgeg eagtteggea 1320
geatetteeg cacetteeac aaccecacet actteteaag gegeetegtg egettetetg 1380
acctetacat ggcetecete agetgeetge teaactaceg egtggaette acettetace 1440
cacgecgtac geogetgeag cacgaggeac cectetggat ggaccagete tgcacegget 1500
geatgaagac cocetteett ggtgacatgg cecacateeg etgagggeac etttattgte 1560
tgggacagge ceteageece tectgeeca tecacecaga caagcaataa aagtggtete 1620
ctccctgaaa aaaaaaaaaa aa
<210> 109
<211> 1818
<212> DNA
<213> Homo sapiens
<220>
<221> misc-feature
<223> Incyte ID No.: 3993377CB1
<400> 100
gaagactete tggageeett caactetetg geaccagage cagtgagtgg aggactatat 60
ggtattgatg acacggaget gatgggtgca gaggacaage tgcctcttga ggacagecet 120
gtgattgetg coettgattg coettcetc aataatgeta etgeetteag teteetggea 180
gatgatagtc aaacatcaac ctctatcttt gccagtccca cctctccacc tgtcctaggg 240
```

90/91

A State of the second of the s

WO 00/44900 PCT/US00/02237

```
gagtetates tacaagataa cagetttaas etgaataata atagtgacge taaacaggaa 300
gaaatggaaa ctcaatcttc agacttccca ccatccctga cccagccagc tcctgatcaq 360
tcatccacta ttcagctaca tccagcaacc tcaccagcag tctcgccaac aacctcccca 420
quagtotece tagtggttte tecageagee tececagaaa tetetecaga agtttgteec 480
gragetteta cagttqtete tecageagte tteteagtgg tetetecage tteeteagea 540
gtecteccag cagteteett agaagteeeg ttgacggett cagtgacate cecaaaagee 600
tetecegtaa etteceeage agetgeettt eeaacageet eeecagcaaa taaggatgte 660
agcagettte tagaaaccae tgetgaegtg gaagagatea etggagaagg acteaetget 720
totggtagtg gtgatgtcat gaggagacgt attgctaccc cagaagaagt togtottccc 780
ctccaacatg ggtggcggag agaggtgcgc atcaagaaca gcagccaccg atggcagggg 840
gagacetggt attatggccc etgtgggaag aggatgaagc aatttccaga agtgatcaag 900
tacetgagee geaacgtggt acacagtgte egeegagage actteagett eagteceegt 960
atgeetgttg gagatttett tgaagaaaga gacaegeeag agggettgea gtgggtgeag 1020
ctctcagcag aggagatccc gtcgaggatt caggcaatta ctggcaaacg gggtcgacct 1080
cgaaacactg agaaggctaa gactaaggaa gtccccaagg tgaaacgggg tcqagqtcqq 1140
ccacctaagg tcaaaatcac tgagctattg aacaagacag acaaccgcc cctaaagaaa 1200
ctggaggccc aagaaacatt gaatgaggag gataaagcaa agattgctaa aagcaagaag 1260
aagatgagge agaaggttea acggggagag tqtcagacta ctatccaagg gcaggccaga 1320
aataagegga aacaagagac caagagetta aageagaagg aagetaagaa gaaatecaag 1380
gctgagaaag aaaaaggaaa gacaaagcag gaaaaactga aggaaaaagt caagagggaa 1440
aagaaggaga aggtaaaaat gaaggaaaag gaggaggtga ccaaagccaa gccagcctgt 1500
aaagcagata agaccetgge cacacagagg cgettggagg aacggcagag gcagcagatg 1560
atcttggagg acatgaagaa gccgacagag gatatgtgtc tgactgacca ccagccctg 1620 cctgacttct cacgagtccc tggtctgaca ttgcccagtg gagccttctc agactgcttg 1680
accattgtgg agttectgea tagetttgge aaggtgetgg geettgatee tgeecaaggt 1740
tgtgcctage ctggggggtc ctgcaggaag gggctcctgt gttcaaggtg accaetctqq 1800
gtgaaggtgc aaaacccg
<210> 110
<211> 785
<212> DNA
<213> Homo sapiens
-220×
<221> unsure
<222> 738
<223> a or g or c or t, unknown, or other
<223> Incyte ID No.: 4717936CB1
<400> 110
gtetetqtag teatgagget gaagggggtg gggacagtgt tgataaaagg cactagagge 60
ageteccaca ecetteeteg gaactgttge ccacatgeag ecceggacae ageceetage 120
ccaaaccta ccettettee teggaggge ccetegagae actgggetge gggtgeetgt 180
cattaagatg ggcacagggt gggagggctt ccagcggacc ctgaaggaag tcgcctacat 240
cetectetge tgetggtgta teaaggaact getggattaa tggtageagg gaactgeete 300
ctctccccac cagcaccatg gctggcatcg ctcaggtggg cagggtagag taaacaggag 360
gcatagetge agettetgtg geagagettg cettagette teattetett etttagecce 420
cagoccaatt gccatcaaga ctootgaago cagotgtgot tgaccaagga tgggccataa 480
acaatgagta aacagtaaag tgtggatcct gctttgagct gtgtcatcta gcagacctgc 540
ctcatctctq agectecttc atteccaacc cetgeetetg gtggacetet ggtgggcagg 600
agettqqact qetetqqeta ggaccecagt aagattgtgg caagacetgg cactceteca 660
agettggcae agtgagecea ceageteaga tggttgatet taccatacce teatagtace 720
aaqaatqqac tqcccccnta agaaacctgt ttgaqaatca ctgaattata aatataaacc 780
```

in which the fall the way were

cagat